Table 1. Surgical procedures used to treat epicondylitis^a

Category	Type of surgery
Denervation	Complete denervation
	Partial lateral denervation
	Partial ventral denervation
Nerve decompression	Decompression of thePIN
	Decompression of the radial nerve
	Combination of denervation and decompression of the PIN
Lengthening of the ERCB	Distal lengthening of the ECRB
	Proximal lengthening of the ERCB
Removal of tissues	Incision of the ERCB
	Partial resection of the annular ligament (Bosworth technique)
	Epicondylar osteotomy
	Epicondylectomy and excision of the distal portion of the annular ligament
	Excision of subtendinious pathological tissue
	Excision of the subcutaneous tissue
	Excision of the radiohumoral bursa
	Fasciectomy of the common extensor origin
	Fasciectomy plus anconeous transfer
	Debriding of the elbow join

^a Adapted from Wilhem et al.⁸⁴ PIN = posterior interosseus nerve ERCB = extensor carpi radialis brevis tendon

De Quervain's Disease

Signs and Symptoms

De Quervain's disease is characterized by pain localized on the radial border of the wrist that may also radiate into the thumb and forearm. The pain is usually worsened by abduction and/or extension of the thumb. Other symptoms may include weakness of the thumb and loss of grip. Range of motion of the wrist and thumb is usually unaffected or only slightly limited.

Table 2. Number of articles Included for Each Key Question

Question #	Carpal Tunnel	Cubital Tunnel	Epicondylitis	De Quervain's
1	189	20	10	0
2	145	32	19	3
3	44	3	50	1
4	12	11	3	1
5	5	14	7	1
6	21	15	6	1
8	0	0	0	0
9	8	0	3	0
10	2	0	2	0
11	12	0	0	0

For the two questions that were not condition specific, Questions 12 and 13, we included 0 and 2 articles, respectively. Question 7 is not depicted in the above table because we addressed it using information from a national database, not published articles.

Evaluating Literature Quality

Because this is a "best evidence" synthesis, we incorporated studies that represented the best available evidence, not the best possible evidence. Therefore, not all evidence that we included is of equal quality.

The quality of studies of treatments that we evaluated can be ranked according to the following hierarchy:

Randomized controlled trials

Other prospective controlled trials

Retrospective controlled trials, including those with historical control groups

Prospective case series

Retrospective case series

Table 3. Coding of Patient Inclusion Criteria

Code	Definition	
WRUED groups		
Symptoms/presented	Patients had unspecified symptoms of the disorder being studied, or were referred for diagnosis of suspected WRUED	
Simple signs/symptoms	Patients included if they had specified symptoms of the disorder, but other tests such as nerve conduction tests were not used for patient selection	
Simple NCS	Patients included if they had abnormal results in a specific nerve conduction test or tests (no more than three tests in selection algorithm)	
Complex objective standard	A specified algorithm with more than three nerve conduction studies or combining specific NCS tests with specific symptoms	
Unspecified (diagnosed)	Authors reported that all patients had been diagnosed with the disorder in question, but did not detail how the diagnosis was defined	
Other	Details reported in separate database field	
Control groups		
Healthy volunteers	Subjects drawn from hospital or community populations, and not being evaluated for other upper extremity disorders	
Workers at risk	Asymptomatic individuals considered to be at risk for WRUED	
Unrelated disease	Subjects were being evaluated or treated for known abnormalities of the hand or wrist unrelated to WRUEDs	
Contralateral arm	Unaffected contralateral extremity of persons with diagnosed WRUED	
Other	Details reported in separate database field	

Table 4. Coding of Diagnostic Test Groups

Test group	Included tests	
Imaging tests	Radiography (film x-ray), computed tomography, MRI, ultrasound	
Nerve conduction	Amplitude, latency, and velocity of signal conduction in median and ulnar nerves	
Composite nerve conduction	Differences and ratios of nerve conduction test results	
Signs and symptoms	Phalen's maneuver, reverse Phalen maneuver, Tinel's sign, Durkin (carpal compression) test, sensory diagrams	
Sensory tests	Semmes-Weinstein monofilament test, vibrometry, current perception threshold	

Table 5. Coding of Results Reporting Level

Reporting level	Definition
Patient-level	Results for each patient reported individually. This includes studies where patient-level results were reported in a graph rather than a table. Where possible, ECRI research analysts
Counts	Sufficient data to yield a two-by-two truth table relating test results to another condition (usually patient's assignment to disease or control group)
Summary statistics	Mean and standard deviation of results for all patients in the group
Agreement or difference	Statistics reporting agreement or difference between results of one test and another, but not the results themselves
Technical criteria	Accuracy, precision, and reproducibility of the test results, but not the results themselves.

Table 6. Coding of Studies of Special Interest

Characteristic	Definition
Longitudinal data	Study reported repeated measurements on the same subjects, from which information on the progression of the condition can possibly be derived
Early diagnosis	Study reported that it was intended to identify early-stage disease. For purposes of this assessment, we relied on the authors' own definitions of "early diagnosis" and did not try to validate that validate that description.
Screening study	Study included at least one group of subjects that can be considered a screening population (e.g. asymptomatic individuals whose work entails repetitive movements).

Peer Review

To select peer-reviewers for the draft evidence report, ECRI prepared a list of 30 potential reviewers. This list was submitted to AHRQ, which approved all reviewers. Letters inviting these individuals to review were then mailed. Fifteen individuals responded to these letters, 12 individuals agreed to review the draft evidence report, and 9 individuals returned reviews.

Upon receipt of reviews, ECRI revised the draft report accordingly. ECRI also prepared a document describing the disposition of all substantive reviewer comments and supplied this document to AHRQ for review and approval.

Table 7. Clinical Signs and Symptoms Used to Diagnose CTS

Test	Definition
Closed fist test ¹⁰¹	The patient makes a fist. If the patient feels tingling within one minute, the test is positive.
Combined Phalen's and Durkan's test ¹⁰²	With the patient's elbow extended, the forearm in supination, and the wrist flexed to 60 degrees, the examiner uses one thumb to apply pressure over the carpal tunnel. If the patient feels tingling or numbness within 30 seconds, the test is positive.
Decreased muscle strength 103	Maximum force exerted by the patient on a measurement device.
Durkan compression test ¹⁰⁴	This test is also called the carpal compression test. With the patient's wrist in a neutral position and the forearm supinated, the examiner uses his/her thumbs to compress the wrist at the median nerve. If the patient feels numbness or tingling within 30 seconds, the test is positive.
Flick test ¹⁰⁵	The patient is asked: "What do you do with your hands when your symptoms are at their worst?" If the patient shakes or flicks the hands, the test is positive.
Gilliat tourniquet test ¹⁰⁶	The examiner inflates a blood pressure monitor on the patient's arm proximal to the elbow. If the patient feels numbness or tingling within one minute, the test is positive.
Grip strength 107	Force measured when patient squeezes a measurement device using the whole hand.
Hypesthesia ¹⁰³	Also called hypoesthesia. It refers to decreased sensitivity to touch.
Pain on VAS ¹⁰⁸	Pain as measured by a visual analog scale in which the patient rates the subjective degree of pain by placing a mark on a graphical bar.
Paresthesia in APB ¹⁰⁹	Tingling in the abductor pollicus brevis muscle of the hand.
Phalen's test ⁸	This test is also called the wrist flexion test. The patient places both elbows on a horizontal surface with the forearms vertical, and allows the wrists to flex by gravity. If the patient feels numbness or tingling within one minute, the test is positive.
Pinch strength ¹⁰⁷	Force measured when patient squeezes a measurement device using the thumb and a finger
Symptoms measured systematically ²⁹	Any symptoms of carpal tunnel such as pain, tingling, or numbness, as measured by a questionnaire or a hand diagram.
Symptoms during ultrasound ¹¹⁰	Whether the patient experiences carpal tunnel symptoms when the wrist is stimulated with an ultrasound transducer.
Reverse Phalen's test ¹¹¹	This test is also called the wrist extension test. The patient extends both wrists and fingers. If the patient feels numbness or tingling within two minutes, the test is positive.
Thenar atrophy ¹⁰³	The degree of wasting in the thenar muscle of the hand.
Thenar weakness ³¹	The degree of weakness in the thenar muscle of the hand.
Tinel's test ⁹	This test is also called Hoffman-Tinel's test. The examiner taps lightly on the medial aspect of the wrist. If the patient feels tingling, the test is positive.

Sources: Massy-Westrop¹¹² and ECRI review of clinical trial articles

Table 8. Sensory tests for Diagnosis of CTS

Test	Definition
Current perception ¹¹³	Whether the patient's threshold for perception of electrical current is within normal limits.
Moving two-point discrimination ¹⁰⁷	The examiner touches two closely-spaced prongs to patient's fingers and moves them distally. The test is positive if the patient cannot discriminate the prongs when they are 4-6 millimeters apart.
Object identification ¹¹⁴	The patient blindly feels wooden shapes and is asked to identify them.
Pinprick sensation ¹⁰⁹	Whether the patient has normal pinprick-induced sensation.
Pressure measurement ¹¹⁵	Whether the patient's threshold for perception of pressure is within normal limits.
Ridge threshold ¹¹⁶	The patient places an index finger on a circular disc that has a small ridge. If the patient's threshold for detection of the ridge is abnormal, the test is positive.
Semmes-Weinstein monofilamen ^{go}	This test is also called the von Frey hairs test. The examiner touches the patient with a series of standardized nylon monofilaments, and records the smallest monofilament the patient can detect the presence of.
Static two-point discrimination ³¹	The examiner touches two closely-spaced prongs to patient's fingers and holds them still. The test is positive if the patient cannot discriminate the prongs when they are 5 millimeters apart.
Temperature measurement ¹¹⁷	Whether the patient's threshold for perception of temperature, heat pain or cold pain is within normal limits.
Tuning fork ³⁰	The examiner hits a metal tuning fork which vibrates, and the patient's threshold for detection of vibration is determined. If the threshold falls outside of normal limits, the test is positive.
Vibrometer ¹¹⁸	An instrument vibrates at varying frequencies, and the patient's threshold for detection of vibration is determined. If the threshold falls outside of normal limits, the test is positive

Sources: Massy-Westrop¹¹² and ECRI review of clinical trial articles

Table 9. Definitions of Nerve Conduction Parameters

Test	Definition
	Nerves tested
Median nerve	The central nerve that is believed to be impaired in carpal tunnel syndrome. It innervates the thumb, index, middle, and ring fingers.
Ulnar nerve	The nerve on the medial side of the arm that innervates the ring and little fingers. Some researchers compare median and ulnar nerve conduction tests to diagnose carpal tunnel syndrome.
Radial nerve	The nerve on the lateral side of the arm that innervates the thumb. Some researchers compare median and radial nerve conduction tests to diagnose CTS.
Motor or sensory	Whether the test assesses motor or sensory nerve function.
Orthodromic or antidromic	The relative placement of the stimulating and recording electrodes. If the stimulating electrode is distal to the recording electrode (i.e., the stimulator is further from the torso), the test is orthodromic. Conversely, if the stimulating electrode is proximal to the recording electrode, (i.e., the stimulator is closer to the torso), the test is antidromic. These terms apply to sensory tests but not to motor tests.
	Electrode placement sites
Abductor pollicus brevis muscle (APB)	A muscle in the hand that is used to record median motor parameters.
Abductor digiti minimi (ADM)	A muscle in the hand that is used to record ulnar motor parameters.
	Parameters Measured
Latency	The time in milliseconds (ms) between stimulation and recording of an electrical impulse.
Onset latency	The time in milliseconds (ms) between stimulation and recording of an electrical impulse when measured to the beginning of the action potential.
Peak latency	The time in milliseconds (ms) between stimulation and recording of an electrical impulse when measured to the largest amplitude of the action potential.
Velocity	Speed of nerve conduction in meters per second (m/s)
Amplitude	Size of the action potential in microvolts (uV)
Presence/absence	Whether the nerve action potential was recordable. In severe cases, some action potentials may not be recordable.
Inching test	A series of nerve conduction tests designed to locate specific areas of nerve slowing. It can be performed orthodromically or antidromically. Electrodes are placed in 9-12 locations which are each a small distance (e.g., 1 cm) apart. By stimulating a fixed site (e.g., the middle finger) and recording at several locations (e.g., 9 evenly-spaced locations along the wrist), researchers can measure the nerve latencies and velocities for each segment along the nerve.

Table 10. Imaging Modalities for the Diagnosis of CTS

Test	Definition
Film	Plain film radiograph (x-ray).
СТ	Computed tomography scan. No articles reported use of obsolete (first or second-generation CT scanners).
MRI	Magnetic resonance imaging scan. No articles reported use of obsolete or prototype MR scanners
Ultrasound	Ultrasonic imaging

Evidence Base

Articles were included in this analysis if they reported counts of positive and negative test results for at least one test, and they included ten or more patients. Having sufficient data from each included study to complete the 2 x 2 diagnostic truth table is important, because sensitivity and specificity must be measured simultaneously, using the same diagnostic threshold. Otherwise, the threshold could be shifted to favor the reported statistic at the expense of the unreported one.

Not all of the articles we examined are addressed in this evidence report. However, data from the articles we did not address are provided in the evidence tables in the appendix. We included articles in these evidence tables, regardless of their level of reporting, if their authors described them as screening studies or studies on "early diagnosis" of CTS.

The evidence tables thus list 205 articles that met our *a priori* inclusion criteria. We subsequently excluded 16 of them. Each of these excluded articles is listed in Table 11 along with its reason for exclusion. Some articles were excluded for more than one reason, but only the first reason is listed in the table. Therefore, this table cannot be used to determine what percentage of the literature suffered a specific flaw. The reasons for exclusion of each study in the table were each confirmed by a second analyst. In case of disagreement, the study was not excluded.

After these exclusions, 189 articles remained for analysis, with a total of 38,087 participants in these studies. The majority of studies (110 or 58%) were conducted outside the United States, and almost all of the studies (184 or 97%) were done at a single center.

In order to be included in meta-analyses of diagnostic trial results, articles had to report sufficient data to permit calculation of sensitivity and specificity for the test in question. In other words, counts of positive and negative test results had to be reported, percentages had to be reported with sufficient data on numbers of patients and controls for us to recalculate the 2 x 2 table, or results for each individual patient had to be reported. Patient-level data were reported in 19 of the 189 articles, and counts for at least some patient groups were reported in 131. Only summary statistics (typically group means) were reported in 39 articles. Even though sensitivity and specificity were not reported in

these articles, they were included in the analysis because they met other criteria, such as reporting "early diagnosis" of CTS or an intent to evaluate diagnostic tests in a screening population. In 129 of the articles (68%), it was possible to determine sensitivity and specificity for at least one test from the reported data; in 79 of the articles, the authors themselves reported sensitivity and specificity.

Table 11. Excluded Studies

Author	Reason for Exclusion	
Ikegaya ¹¹⁹	Special patient population (dialysis)	
Tackmann ¹²⁰	No diagnostic data	
Jordan ¹²¹	Reported only statistical significance of results	
Sivri ¹²²	Special patient population (arthritis), only 2 cases of CTS	
Sto Ip-Smith ¹²³	Special patient population (pregnant women), only 5 cases of CTS	
Dlabalová ¹²⁴	All patients post-surgery for CTS	
Lazaro ¹²⁵	All patients post-surgery for CTS	
Nakamichi ¹²⁶	All patients post-surgery for CTS	
Williams ¹²⁷	Discrepancies in reported results; 2 x 2 table could not be accurately reproduced by ECRI.	
Mossman ¹²⁸	Published as letter rather than full paper; 2 x 2 table could not be accurately reproduced by ECRI.	
Westerman ¹²⁹	Discrepancies in reported number of patients, unexplained exclusions of patients.	
Herrick ¹³⁰	Combined results from CTS patients and patients with other conditions.	
MacDermid ¹³¹	Combined results from CTS patients and patients with other conditions.	
Gerrning ¹³²	Combined results from CTS patients and patients with other conditions.	
Byl ¹³³	Combined results from CTS patients and patients with other conditions.	
Palmer ¹³⁴	Combined results from CTS patients and patients with other conditions.	

Internal Validity of Results

To evaluate the quality of this literature base, we determined what proportion of articles reported various details of study methods or results. Reporting of these details is necessary to verify the internal validity and generalizability of study results. Reporting of characteristics affecting the internal validity of the results (the degree to which the reported results reflect the true performance of the test in the conditions of the particular study) is summarized in Table 12; this table includes all 189 articles on CTS diagnosis that were abstracted into the database. Details of the studies eventually included in quantitative analyses are listed in Table 13.

The design of most studies raised the possibility of age bias in which patients were markedly older than controls. Some nerve conduction measurements become slower as people age, ⁹⁷ thus if patients are older than controls, the study will overestimate the

Table 12. Summary of Study Characteristics Affecting Internal Validity

Study characteristic	Number of studies reporting (percentage)	Specifics (percentage)
Whether trial was funded by a for-profit institution	24 (13%)	For-profit funding: 3 (2%) No for-profit funding: 21 (11%)
Was selection of patients prospective or retrospective?	75 (40%)	Prospective: 58 (28%) Retrospective: 17 (9%)
Patient inclusion criteria	185 (98%)	See Table 46
Patient exclusion criteria	87 (46%)	See Table 46
Was sex distribution of patients reported?	131 (69%)	^a Percentage female: 61.5%
Was the percentage of females in the	89 (47%)	Yes: 65 (34%)
patient group within 20 percentage points of the control group?		No, patients were = 20% more female: 21 (11%)
		No, controls were =20% more female: 3 (2%)
Were patient ages reported?	123 (65%)	^a Mean age 48.1 years
Was the mean patient age within 5 years	89 (47%)	Yes: 52 (28%)
of the mean control age?		No, patients were = 5 years older: 36 (19%) No, controls were =5 years older: 1 (1%)
Was duration of patients' condition reported?	18 (10%)	^{a, b} Mean duration 28.1 months
Were patient comorbidities reported?	46 (24%)	NA
Was the test operator blinded?	13 (7%)	Yes: 13 (7%)
Was the test reader blinded?	23 (12%)	Yes: 23 (12%)
Were there multiple test readers?	7 (4%)	2 readers: 4 (2%)
·	, ,	3 readers: 2 (1%)
		4 readers: 1 (1%)
What was the method for multiple test	4 (57% of studies	Independent: 2 (1%)
readers?	reporting multiple	Mean: 1 (1%)
	readers)	Consensus: 1 (1%)
Was the test compared to an	38 (20%)	Yes: 38 (20%)
ndependent reference standard?	, ,	,
Were all patients given the test and the reference standard?	28 (15%)	Yes: 28 (15%)

Key:
NA—not applicable

a Calculated on a per-patient basis (i.e., weighted by number of patients in each study reporting this characteristic)

b Studies reporting median duration 109,136,137 were excluded from calculation.

Table 13. Study Characteristics Affecting Internal Validity of Results

Article	Funded by for- profit institution?	Inclusion cri- teria reported?		Method of diag- nosis reported		Comorbidity reported	^а Реі	Possible sex bias	^a Mean age	Possible age bias	^a Mean duration of condition	Test operator blinded	Test reader blinded	Multiple readers	Method for multiple readers	Independent reference standard	Were patients given both test and reference
			Dista	al Mot	or Latency:	Unspe	cified	Diagno	osi s F	atient	Group)					
Rosén, 1993 ¹³⁸ NR Yes Yes NR						NR	75%	Р	41	No	NR	NR	NR	NR	NR	No	No
Marin, 1983 ¹³⁹	NR	Yes	NR	NR	NR	NR	86%	Р	49	Р	13	NR	NR	NR	NR	No	No
Kimura, 1979 ¹⁴⁰	NR	Yes	Yes	Yes	NR	NR	75%	No	48	No	NR	NR	NR	NR	NR	No	No
Loong, 1972 141	NR	Yes	NR	NR	NR	NR	100%	No	43.7	MNR	12.7	NR	NR	NR	NR	No	No
Plaja, 1971 ¹⁴²	NR	NR	Yes	NR	Retrospective	NR	NR	GNR	NR	MNR	NR	NR	NR	NR	NR	No	No
			Dista	al Mo	tor Latency:	Symp	toms/F	resen	ted Pa	atient (Groups	3					
Murthy, 1999 143	NR	Yes	NR	Yes	NR	NR	NR	GNR	NR	ANR	NR	NR	NR	NR	NR	No	No
Atroshi, 1996 136	No	Yes	NR	NR	Prospective	Yes	69%	No	52	Р	24	NR	NR	NR	NR	No	No
Kuntzer, 1994 144	NR	Yes	Yes	NR	Prospective	NR	80%	Р	51	Р	NR	NR	NR	NR	NR	No	No
Chang, 1991 ¹⁴⁵	NR	Yes	Yes	NR	NR	Yes	79%	GNR	42.3	No	NR	NR	NR	NR	NR	No	No
Cioni, 1989 ¹⁴⁶	NR	Yes	Yes	NR	NR	NR	16%	С	46.4	Р	NR	NR	NR	NR	NR	No	No
Messina, 1980 120	NR	Yes	NR	NR	NR	NR	NR	GNR	45.1	No	NR	NR	NR	NR	NR	No	No
Melvin, 1972 ¹⁴⁷	NR	Yes	NR	NR	NR	NR	NR	GNR	NR	ANR	NR	NR	NR	NR	NR	No	No
Loong, 1971 ¹⁴⁸	NR	Yes	Yes	NR	NR	Yes	100%	No	NR	ANR	7.6	NR	NR	NR	NR	No	No
		Р	alma	r Sen	sory Latency	: Syn	ptoms	/Prese	ented	Patien	t Grou	ps					
Murthy, 1999 143	NR	Yes	NR	Yes	NR	NR	NR	GNR	NR	ANR	NR	NR	NR	NR	NR	No	No
Girlanda, 1998 ¹⁴⁹	NR	Yes	Yes	NR	NR	Yes	93%	GNR	39	ANR	48	NR	NR	NR	NR	No	No
Chang, 1991 145	NR	Yes	Yes	NR	NR	Yes	79%	GNR	42.3	No	NR	NR	NR	NR	NR	No	No
Jackson, 1989 ¹⁵⁰	No	Yes	Yes	NR	NR	Yes	82%	No	52.6	Р	NR	NR	NR	NR	NR	No	No
Escobar, 1985 151	NR	Yes	Yes	NR	NR	Yes	70%	No	NR	ANR	NR	NR	NR	NR	NR	No	No

Article	Funded by for- profit institution?	Inclusion cri- teria reported?	Exclusion criteria reported	Method of diagnosis reported	Patient selection	Comorbidity	Percent female	Possible sex bias	^a Mean age	Possible age bias	^a Mean duration of condition	Test operator blinded	Test reader blinded	Multiple readers	Method for multiple readers	Independent reference standard	Were patients given both test and reference
	Ē	_ te	Ш Ţ	Šč				_			J _e	_		Ĕ	Me	_	y gi
					Phalen's Ma						,					1	
Szabo, 1999 152	No	Yes	NR	Yes	Prospective	NR	76%	No	NR	ANR	NR	NR	Yes	NR	NR	No	No
Fertl, 1998 153	NR	Yes	Yes	Yes	Prospective	NR	83%	Р	55.5	Р	NR	Yes	Yes	NR	NR	No	No
Gerr, 1998 ³¹	NR	Yes	Yes	Yes	NR	NR	72%	No	46.6	Р	NR	NR	NR	NR	NR	Yes	No
Ghavanini, 1998 ¹⁵⁴	NR	Yes	Yes	Yes	Prospective	NR	81%	No	40	No	15	NR	NR	NR	NR	No	No
Tetro, 1998 ¹⁰²	No	Yes	Yes	Yes	Prospective	NR	64%	No	49.3	No	NR	NR	NR	NR	NR	No	No
González del Pino, 1997 104	NR	Yes	NR	Yes	Prospective	NR	81%	No	50	No	37.9	NR	NR	3	NR	Yes	Yes
De Smet, 1995 101	NR	Yes	NR	Yes	NR	NR	88%	С	49.2	С	NR	NR	NR	NR	NR	No	No
Werner, 1994 111	NR	Yes	NR	Yes	NR	NR	NR	GNR	NR	ANR	NR	NR	NR	NR	NR	No	No
Durkan, 1991 155	No	Yes	NR	Yes	NR	NR	NR	GNR	45	ANR	NR	NR	NR	NR	NR	Yes	Yes
Gellman, 1986 106	No	Yes	NR	Yes	NR	Yes	74%	GNR	NR	ANR	NR	NR	NR	NR	NR	Yes	Yes
					Tinel's S	ign: /	All Pati	ent Gr	oups								
Szabo, 1999 152	No	Yes	NR	Yes	Prospective	NR	76%	No	NR	ANR	NR	NR	Yes	NR	NR	No	No
Gerr, 1998 ³¹	NR	Yes	Yes	Yes	NR	NR	72%	No	46.6	Р	NR	NR	NR	NR	NR	Yes	No
Ghavanini, 1998 154	NR	Yes	Yes	Yes	Prospective	NR	81%	No	40	No	15	NR	NR	NR	NR	No	No
Tetro, 1998 ¹⁰²	No	Yes	Yes	Yes	Prospective	NR	64%	No	49.3	No	NR	NR	NR	NR	NR	No	No
González del Pino, 1997 104	NR	Yes	NR	Yes	Prospective	NR	81%	No	50	No	37.9	NR	NR	3	NR	Yes	Yes
De Smet, 1995 101	NR	Yes	NR	Yes	NR	NR	88%	С	49.2	С	NR	NR	NR	NR	NR	No	No
Durkan, 1991 ¹⁵⁵	No	Yes	NR	Yes	NR	NR	74%	GNR	45	ANR	NR	NR	NR	NR	NR	Yes	Yes
Seror, 1987 ¹⁵⁶	NR	Yes	Yes	Yes	NR	NR	79%	No	56.8	No	NR	NR	NR	NR	NR	No	No
Gellman, 1986 106	No	Yes	NR	Yes	NR	Yes	NR	GNR	NR	ANR	NR	NR	NR	NR	NR	Yes	Yes
Gelmers, 1979 ²⁹	NR	Yes	Yes	Yes	NR	NR	81%	No	57	No	NR	NR	NR	NR	NR	Yes	No
Stewart, 1978 157	NR	Yes	Yes	Yes	NR	Yes	81%	No	55	No	NR	NR	NR	NR	NR	Yes	No

Method for multiple test readers: Indep—Independent

<u>Key:</u>

^aPercent female, mean age, and mean duration of condition for CTS patients

Possible sex bias: No—proportion women in epicondylitis group within 20% of proportion of women in control group; P—Patients were more likely to be female; C—Controls were more likely to be female; GNR—Genders not reported for both groups; NC—Study did not contain a separate control group in the control group within 5 years of mean age of control group; P—Patients were older than controls; C—Controls were older than control controls; C—Controls were older than controls; C—Controls were older than controls; C—Controls were older than controls; C—Controls Possible age bias: No—mean age of epicondylitis group within 5 years of mean age of control group; P—Patients were older than controls; C—Controls were older than patients;

ANR—Ages not reported for both groups; NC—Study did not contain a separate control group

Table 14. Summary of Study Characteristics Affecting Generalizability

Study characteristic	Number of studies reporting (percentage)	Specifics (percentage)
Years in which study was conducted	39 (21%)	NA
Number of centers	189 (100%)	Single: 184 (97%) Multiple (<5): 4 (2%) Multiple (>5): 1 (1%)
Country in which study was conducted	189 (100%)	USA: 79 (42%) Other: 110 (58%)
Patient inclusion criteria	185 (98%)	See Table 46
Patient exclusion criteria	87 (46%)	See Table 46
Were patient comorbidities reported?	46 (24%)	NA
Was sex distribution of patients reported?	131 (69%)	^a Percentage female: 61.5%
Were patient ages reported?	123 (65%)	^a Mean age 48.1 years
Was duration of patients' condition reported?	18 (10%)	^{a, b} Mean duration 28.1 months
Did all patients have previous conservative treatment?	1 (1%)	Yes: 1 (1%)
Did any patients have previous surgical treatment?	6 (3%)	Yes: 6 (3%)
Adequate reporting of study's source of patients	29 (15%)	NA
Was there a potential selection bias for easy cases?	58 (31%)	Yes: 58 (31%)
Was there a potential selection bias for hard cases?	40 (21%)	Yes: 40 (21%)

Key:
NA—not applicable

a Calculated on a per-patient basis (i.e., weighted by number of patients in each study reporting this characteristic)

b Studies reporting median duration 109,136,137 excluded from calculation

Table 15. Study Characteristics Affecting Generalizability of Results

Article	Years in which trial was conducted	Number of centers	Country where trial was conducted	Are patient comorbidity reported?	Percent female	Mean age	Mean duration of condition	Did all patients have previous conservative	Did any patients have previous surgical	Source of patients adequately described and generalizable to broader clinical	Potential selection bias for easy cases?	Potential selection bias for difficult cases?
D 1002 120	100/ 1007	1	Motor Latenc				osis Patie			\/	NI-	NI-
Rosén, 1993 ¹³⁸ Marin, 1983 ¹³⁹	1986-1987 NR	Single Single	Sweden USA	No No	75% 86%	41 49	NR 13	No No	No No	Yes No	No Yes	No No
Kimura, 1979 ¹⁴⁰	1978	Single	USA	No	75%	48	NR	No	No	No	No	Yes
Loong, 1972 ¹⁴¹	NR	Single	Singapore	No	100%	43.7	12.7	No	No	No	No	No
Plaja, 1971 ¹⁴²	NR	Single	Spain	No	NR	NR	NR	No	No	No	Yes	No
Distal Motor Latency: Symptoms/Presented Patient Groups												
Murthy, 1999 143	NR	Single	India	No	NR	NR	NR	No	No	No	Yes	No
Atroshi, 1996 136	NR	Single	Sweden	Yes	69%	52	24	Yes	No	No	Yes	No
Kuntzer, 1994 144	NR	Single	Switzerland	No	80%	51	NR	No	No	Yes	No	No
Chang, 1991 145	NR	Single	Taiwan	Yes	79%	42.3	NR	No	No	No	No	No
Cioni, 1989 146	NR	Single	Italy	No	16%	46.4	NR	No	No	No	No	No
Messina, 1980 120	NR	Single	Italy	No	NR	45.1	NR	No	No	No	No	No
Melvin, 1972 147	NR	Single	USA	No	NR	NR	NR	No	No	No	No	No
Loong, 1971 148	NR	Single	Singapore	Yes	100%	NR	7.6	No	No	No	No	No
			ensory Later	ncy: Syn					-			
Murthy, 1999 143	NR	Single	India	No	NR	NR	NR	No	No	No	Yes	No
Girlanda, 1998 149	NR	Single	Italy	Yes	93%	39	48	No	No	No	No	Yes
Chang, 1991 145	NR	Single	Taiwan	Yes	79%	42.3	NR	No	No	No	No	No
Jackson, 1989 150	NR	Single	Canada	Yes	82%	52.6	NR	No	No	No	No	No
Escobar, 1985 ¹⁵¹	NR	Single	USA	Yes	70%	NR	NR	No	No	No	No	No

	Years in which trial was conducted	Number of centers	Country where trial was conducted	Are patient comorbidity reported?	Percent female	Mean age	Mean duration of condition	Did all patients have previous conservative	Did any patients have previous surgical	Source of patients adequately described and generalizable to	Potential selection bias for easy cases?	Potential selection bias for difficult cases?		
Article	¥ ,		ပိ		Pe		Me	Dic ha	Did	Soul de gel gel	sele	sele		
			Phalen's	Maneuve	r: All	Patient	Groups							
Szabo, 1999 152														
Fertl, 1998 153	1997	Single	Austria	No	83%	55.5	NR	No	No	Yes	No	No		
Gerr, 1998 ³¹	NR	Single	USA	No	72%	46.6	NR	No	No	No	No	No		
Ghavanini, 1998 ¹⁵⁴	NR	Single	Iran	No	81%	40	15	No	No	No	No	No		
Tetro, 1998 ¹⁰²	1995-1997	Single	USA	No	64%	49.3	NR	No	No	Yes	No	No		
González del Pino, 1997 104	1992-1995	Single	Spain	No	81%	50	37.9	No	No	No	Yes	No		
De Smet, 1995 101	NR	Single	Belgium	No	88%	49.2	NR	No	No	No	No	No		
Werner, 1994 111	NR	Single	USA	No	NR	NR	NR	No	No	No	No	No		
Durkan, 1991 ¹⁵⁵	1987-1990	Single	USA	No	NR	45	NR	No	No	No	No	No		
Gellman, 1986 ¹⁰⁶	1982-1984	Single	USA	Yes	74%	NR	NR	No	No	No	Yes	No		
		1				ent Gr		Г						
Szabo, 1999 ¹⁵²	1993-1996	Single	USA	No	76%	NR	NR	No	No	No	No	No		
Gerr, 1998 ³¹	NR	Single	USA	No	72%	46.6	NR	No	No	No	No	No		
Ghavanini, 1998 154	NR	Single	Iran	No	81%	40	15	No	No	No	No	No		
Tetro, 1998 ¹⁰²	1995-1997	Single	USA	No	64%	49.3	NR	No	No	Yes	No	No		
González del Pino, 1997 104	1992-1995	Single	Spain	No	81%	50	37.9	No	No	No	Yes	No		
De Smet, 1995 101	NR	Single	Belgium	No	88%	49.2	NR	No	No	No	No	No		
Durkan, 1991 ¹⁵⁵	1987-1990	Single	USA	No	74%	45	NR	No	No	No	No	No		
Seror, 1987 ¹⁵⁶	NR	Single	France	No	79%	56.8	NR	No	No	No	No	No		
Gellman, 1986 106	1982-1984	Single	USA	Yes	NR	NR	NR	No	No	No	Yes	No		
Gelmers, 1979 ²⁹	NR	Single	Netherlands	No	81%	57	NR	No	No	No	Yes	No		
Stewart, 1978 157	NR	Single	Canada	Yes	81%	55	NR	No	No	No	Yes	No		

Key: NR—not reported

Table 16. Articles Self-Described as "Early Diagnosis" of CTS

Article	Patient selection criteria relevant to early detection	Symptoms and normal NCS?	Authors' proposed method for early detection	Sensor y NCS?
Seror, 2000 ¹⁵⁸	Symptoms, but normal needle examination, normal DML (<4 ms) and normal palm-to-wrist orthodromic SCV (>45 m/s).	Ø	Orthodromic sensory inching test from the middle finger.	☑
Girlanda, 1998 ¹⁴⁹	Symptoms, but no weakness, no muscle atrophy, and normal DML (<4 ms).		Combination of nerve conduction tests:a) Difference between median and ulnar orthodromic SCV from ring finger to wrist, and b) Ratio of orthodromic SCV from middle finger to palm and orthodromic SCV from palm to wrist	V
Seror, 1998 ¹⁵⁹	Symptoms, but normal DML (<4 ms) and normal palm-to-wrist orthodromic SCV (>45 m/s).	₫	Orthodromic sensory inching test from the middle finger.	V
Terzis, 1998 ¹⁶²	Symptoms, but normal DML (<4.2 ms)	Ĭ	Combination of orthodromic sensory nerve conduction tests from the ring finger.	V
Bronson, 1997 ¹⁶³	Symptoms, but normal DML (<4 ms) and normal needle examination.	₫	Comparison of DMLs using five different wrist positions.	?
Murata, 1996 ¹⁶⁴	Workers at risk	?	Ratio of:a) Antidromic SCV from wrist to index finger, and b) Antidromic SCV from palm to index finger	Ø
Padua, 1996 ¹⁶⁵	Symptoms, but no signs of severe CTS (e.g., absent SNAP at the wrist).	Ĭ	Ratio of:a) Orthodromic SCV from middle finger to palm, and b) Orthodromic SCV from palm to wrist	V
Young, 1995 ¹⁶⁶	Workers at risk	?	Total score on a grading scale that included seven clinical signs, four symptoms, and DML≥4.45 ms.	?
Johnson, 1993 ¹⁶⁷	Workers at risk	?	Track changes in DML over time	?
Uncini, 1993 ¹⁶⁰	Symptoms, but normal DML (<4.2 ms) and normal SCV from index finger to wrist (>45 m/s)	Ø	Difference between: a)Median orthodromic latency between ring finger and wrist, and b) Ulnar orthodromic latency between ring finger and wrist	V
Jetzer, 1991 ¹⁶⁸	Workers at risk	?	Vibrometry	?
Luchetti, 1991 ¹⁶⁹	Symptoms, but normal motor function, sensory function, quantitative sensory examination, cutaneous trophism, DSL (NR), and DML (NR).	V	Antidromic inching test to the middle finger	V

Article	Patient selection criteria relevant to early detection	Symptoms and normal NCS?	Authors' proposed method for early detection	Sensor y NCS?
Charles, 1990 ¹⁷⁰	Clinical diagnosis of CTS by referring physician, and at least one of the following: a) DML≥4.5 ms; b) Orthodromic SCV from index finger <45 m/s; c) Difference ≥0.5 ms between median and ulnar sensory antidromic latencies to the ring finger	?	Difference between: a) Median antidromic latency between ring finger and wrist, and b) Ulnar antidromic latency between ring finger and wrist	Ø
Palliyath, 1990 ¹⁷¹	Symptoms, but "very little electrophysiological changes on routine tests for CTS" (p 307).	V	Duration of relative refractory period and absolute refractory period.	?
Cioni, 1989 ¹⁴⁶	Symptoms	?	Orthodromic SCV from ring finger to wrist	$\overline{\mathbf{A}}$
Jackson, 1989 ¹⁵⁰	Symptoms. Patients were stratified into three groups, and one group represented mild CTS as defined by normal NCS (based on four tests) and normal needle examination.	V	Combination of two nerve conduction tests: a) Difference between median and radial antidromic sensory latencies from wrist to thumb, and b) Difference between median and ulnar antidromic sensory latencies from wrist to ring finger	V
Uncini, 1989 ¹⁶¹	Symptoms, but normal DML (≤4.2 ms) and SNAPs were present with normal amplitude.	V	Difference between:a) Median orthodromic latency between ring finger and wrist, and b) Ulnar orthodromic latency between ring finger and wrist	V
Wongsam, 1983 ¹⁷²	Symptoms suggesting early CTS.	?	Ratio of:a) Antidromic latency from wrist to middle fingerb) Antidromic latency from palm to middle finger	Ø

Key:

DML—Distal motor latency

DSL—Distal sensory latency

ms—Milliseconds

m/s—Meters per second

SCV—Sensory conduction velocity

SNAP—Sensory nerve action potential

NR—Not reported

"Diagnosis Studies"

Our evaluation of methods for diagnosing CTS is primarily meta-analytic. To identify diagnostic tests of CTS for which meta-analyses were appropriate, we performed several tabulations. These tabulations were restricted to studies that met each of the following three criteria: 1) Study included a carpal tunnel syndrome group; 2) Study included a normal group; 3) Study was not a screening study. There were 138 studies that met all of these criteria.

For each test, we determined the number of studies in each of four patient selection categories that reported the test. Within each of these four categories, we also determined the number of studies for which sensitivity and specificity could be derived (based on information provided in the article). These study counts appear in Table 17 through

Table 17. Numbers of Studies Reporting Signs/Symptoms Tests Across Patient Selection Categories

Legend:

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Sign/symptom	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified diagnosis
Closed fist test	0, 0	1, 1	1, 1	0, 0
Combined Phalen's/Durkan test	1, 1	0, 0	0, 0	0, 0
Decreased muscle strength	0, 0	0, 0	1, 1	0, 0
Durkan compression	5, 5	1, 1	3, 3	1, 1
Flick sign	0, 0	0, 0	0, 0	1, 1
Gilliat tourniquet	1, 1	1, 1	1, 1	1, 1
Grip strength	0, 0	0, 0	0, 0	1, 0
Hypesthesia	0, 0	0, 0	1, 1	0, 0
Pain on VAS	0, 0	0, 0	1, 1	1, 1
Paresthesia in APB	0, 0	0, 0	0, 0	1, 1
Phalen's/reverse Phalen's	7, 7	2, 1	6, 6	3, 3
Pinch strength	0, 0	0, 0	0, 0	1, 0
Symptoms measured systematically	3, 3	0, 0	2, 2	1, 0
Symptoms during ultrasound	0, 0	0, 0	1, 1	0, 0
Thenar atrophy	0, 0	0, 0	2, 2	0, 0
Thenar weakness	0, 0	0, 0	1, 1	0, 0
Tinel's	9, 9	2, 1	3, 3	2, 2

Table 18. Numbers of Studies Reporting Sensory Tests Across Patient Selection Categories

Legend:

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Sensory test	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified diagnosis
Object identification	0, 0	0, 0	0, 0	1, 0
Pinprick sensation	0, 0	0, 0	0, 0	1, 1
Pressure measurement	0, 0	0, 0	1, 1	1, 0
Ridge threshold	0, 0	0, 0	0, 0	1, 0
Semmes-Weinstein	1, 1	0, 0	0, 0	4, 1
filament				
Temperature	0, 0	0, 0	1, 1	2, 1
measurement				
Texture discrimination	0, 0	0, 0	0, 0	1, 0
Tuning fork	1, 1	0, 0	1, 1	0, 0
Two-point	2, 2	0, 0	2, 2	1, 0
discrimination (moving				
or static)				
Vibrometer	2, 2	0, 0	5, 5	1, 0

Table 19. Numbers of Studies Reporting Nerve Conduction Tests Across Patient Selection Categories

Legend:

Nerve tested: MED-median, RAD-radial, ULN-ulnar MOT-motor, SEN-Sensory

Configuration (not applicable to motor nerve tests): OR-orthodromic, AN-antidromic Stimulation electrode placement: ELB-elbow, FOR-forearm, WR-wrist, PAL-palm, TH-thumb, IN-index finger, MI-middle finger, RI-ring finger, LI-little finger, APB-abductor policis brevis, ADM-abductor digiti minimi, OTH-other

Recording electrode placement (see D for abbreviations)

Measured parameter: LAT-latency, PRE-presence/absence of signal, AMP-amplitude, VEL-velocity, INCH-inching, OTH-other

Blank cells—Not reported or not applicable

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Shaded cells—Ten or more articles reporting sensitivity and specificity.

	Nerve	Condu	ction Tes	t		Patient selection type					
Nerve Tested	Nerve Tested	Configuration	Stimulation	Recording	Parameter	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified diagnosis		
	MOT				LAT	1, 1	0, 0	0, 0	1, 1		
	MOT		WR	OTH	LAT	0, 0	0, 0	0, 0	1, 1		
	SEN				LAT	1, 1	0, 0	0, 0	1, 1		
	SEN	OR	TH	WR	PRE	0, 0	0, 0	1, 1	0, 0		
MED					OTH	0, 0	0, 0	2, 2	0, 0		
MED	MOT					0, 0	0, 0	0, 0	1, 0		
MED	MOT				AMP	0, 0	1, 0	0, 0	0, 0		
MED	MOT				LAT	2, 1	1, 0	2, 2	2, 1		
MED	MOT				OTH	1, 1	1, 0	2, 1	0, 0		
MED	MOT				VEL	0, 0	1, 0	1, 1	1, 0		
MED	MOT			APB	AMP	1, 1	0, 0	0, 0	0, 0		
MED	MOT			APB	LAT	1, 1	0, 0	0, 0	0, 0		
MED	MOT		ELB	APB	AMP	1, 0	0, 0	1, 1	1, 1		
MED	MOT		ELB	APB	LAT	1, 1	0, 0	0, 0	1, 1		
MED	MOT		ELB	APB	OTH	1, 1	0, 0	1, 1	0, 0		
MED	MOT		ELB	APB	VEL	1, 0	0, 0	1, 1	2, 2		
MED	MOT		ELB	IN	AMP	0, 0	0, 0	0, 0	1, 1		
MED	MOT		ELB	IN	LAT	0, 0	0, 0	0, 0	1, 1		
MED	MOT		ELB	IN	VEL	0, 0	0, 0	0, 0	1, 1		
MED	MOT		ELB	WR	AMP	1, 0	0, 0	0, 0	0, 0		
MED	MOT		ELB	WR	LAT	0, 0	0, 0	0, 0	0, 0		
MED	MOT		ELB	WR	VEL	2, 1	0, 0	3, 3	1, 1		
MED	MOT		FOR		VEL	1, 1	0, 0	0, 0	1, 1		
MED	MOT		FOR	APB	AMP	1, 1	0, 0	0, 0	0, 0		

	Nerve	Condu	Patient selection type						
Nerve Tested	Nerve Tested	Configuration	Stimulation	Recording	Parameter	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified
MED	MOT		FOR	APB	LAT	1, 1	0, 0	0, 0	0, 0
MED	MOT		FOR	APB	VEL	0, 0	0, 0	1, 1	0, 0
MED	MOT		FOR	PAL	AMP	1, 1	0, 0	0, 0	0, 0
MED	MOT		FOR	PAL	LAT	1, 1	0, 0	0, 0	0, 0
MED	MOT		FOR	WR	VEL	0, 0	0, 0	0, 0	1, 1
MED	MOT		PAL	APB	AMP	1, 1	0, 0	1, 0	2, 1
MED	MOT		PAL	APB	LAT	0, 0	0, 0	0, 0	2, 1
MED	MOT		PAL	IN	AMP	0, 0	0, 0	0, 0	1, 1
MED	MOT		PAL	IN	LAT	0, 0	0, 0	0, 0	1, 1
MED	MOT		PAL	IN	VEL	0, 0	0, 0	0, 0	1, 1
MED	MOT		WR		AMP	0, 0	0, 0	1, 1	0, 0
MED	MOT		WR		LAT	2, 2	1, 0	1, 1	0, 0
MED	MOT		WR		PRE	1, 1	0, 0	0, 0	0, 0
MED	MOT		WR		VEL	1, 1	0, 0	0, 0	0, 0
MED	MOT		WR	APB	AMP	2, 1	0, 0	9, 7	9, 6
MED	MOT		WR	APB	LAT	4, 4	3, 2	21, 17	24, 21
MED	MOT		WR	APB	OTH	2, 1	1, 0	1, 1	2, 2
MED	MOT		WR	APB	PRE	0, 0	0, 0	3, 3	1, 1
MED	MOT		WR	APB	VEL	0, 0	0, 0	2, 1	5, 5
MED	MOT		WR	IN	AMP	0, 0	0, 0	0, 0	1, 1
MED	MOT		WR	IN	LAT	0, 0	0, 0	0, 0	1, 1
MED	MOT		WR	IN	VEL	0, 0	0, 0	0, 0	1, 1
MED	MOT		WR	OTH	AMP	1, 0	0, 0	1, 1	1, 1
MED	MOT		WR	OTH	LAT	1, 1	1, 1	8, 8	3, 3
MED	MOT		WR	OTH	OTH	0, 0	0, 0	0, 0	1, 1
MED	MOT		WR	OTH	VEL	1, 0	0, 0	1, 1	0, 0
MED	MOT		WR	PAL	AMP	0, 0	0, 0	1, 1	0, 0
MED	MOT		WR	PAL	LAT	0, 0	0, 0	1, 1	0, 0
MED	MOT		WR	PAL	OTH	0, 0	0, 0	0, 0	1, 1
MED	MOT		WR	PAL	VEL	0, 0	0, 0	1, 0	0, 0
MED	MOT		WR	TH	LAT	0, 0	0, 0	2, 0	0, 0
MED	MOT		WR	TH	VEL	0, 0	0, 0	1, 1	0, 0
MED	SEN				T A.T.	0, 0	0, 0	0, 0	1, 0
MED	SEN				LAT	3, 2	0, 0	0, 0	1, 0
MED	SEN				OTH	0, 0	0, 0	1, 0	1, 0
MED	SEN		MAD		VEL	0, 0	1, 0	0, 0	0, 0
MED	SEN		WR		AMP	1, 1	0, 0	0, 0	0, 0
MED	SEN	A 3.7	WR		LAT	1, 1	0, 0	0, 0	0, 0
MED	SEN	AN			AMP	0, 0	0, 0	0, 0	1, 1
MED	SEN	AN			LAT	1, 1	0, 0	1, 1	1, 1
MED	SEN	AN	ELD	TNI	VEL	1, 1	0, 0	1, 1	0, 0
MED	SEN	AN	ELB	IN	AMP	0, 0	0, 0	1, 1	0, 0
MED	SEN	AN	ELB	IN	OTH	0, 0	0, 0	1, 1	0, 0

	Nerve	Condu	Patient selection type						
Nerve Tested	Nerve Tested	Configuration	Stimulation	Recording	Parameter	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified
MED	SEN	AN	ELB	MI	VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	AN	ELB	PAL	INCH	0, 0	0, 0	0, 0	1, 1
MED	SEN	AN	ELB	WR	VEL	0, 0	0, 0	2, 2	1, 1
MED	SEN	AN	FOR	IN	LAT	0, 0	0, 0	1, 0	0, 0
MED	SEN	AN	FOR	RI	LAT	0, 0	0, 0	1, 0	0, 0
MED	SEN	AN	FOR	TH	LAT	0, 0	0, 0	1, 0	0, 0
MED	SEN	AN	PAL	IN	AMP	1, 1	0, 0	2, 1	0, 0
MED	SEN	AN	PAL	IN	LAT	0, 0	0, 0	1, 1	0, 0
MED	SEN	AN	PAL	IN	PRE	1, 1	0, 0	0, 0	0, 0
MED	SEN	AN	PAL	IN	VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	AN	PAL	MI	, LL	0, 0	0, 0	1, 0	0, 0
MED	SEN	AN	PAL	MI	AMP	0, 0	0, 0	2, 1	0, 0
MED	SEN	AN	PAL	MI	LAT	0, 0	0, 0	2, 1	0, 0
MED	SEN	AN	PAL	MI	OTH	0, 0	0, 0	1, 1	0, 0
MED	SEN	AN	PAL	MI	VEL	0, 0	0, 0	1, 1	2, 2
MED	SEN	AN	WR		LAT	0, 0	0, 0	1, 1	0, 0
MED	SEN	AN	WR	IN	AMP	3, 2	0, 0	6, 5	5, 4
MED	SEN	AN	WR	IN	LAT	1, 1	0, 0	11, 9	5, 3
MED	SEN	AN	WR	IN	OTH	2, 1	0, 0	2, 2	0, 0
MED	SEN	AN	WR	IN	PRE	1, 1	0, 0	2, 2	2, 2
MED	SEN	AN	WR	IN	VEL	0, 0	0, 0	3, 2	0, 0
MED	SEN	AN	WR	MI	AMP	0, 0	0, 0	4, 3	0, 0
MED	SEN	AN	WR	MI	INCH	1, 1	0, 0	1, 1	0, 0
MED	SEN	AN	WR	MI	LAT	0, 0	0, 0	2, 1	0, 0
MED	SEN	AN	WR	MI	PRE	0, 0	0, 0	1, 1	0, 0
MED	SEN	AN	WR	MI	VEL	0, 0	0, 0	3, 3	1, 1
MED	SEN	AN	WR	OTH	VEL	0, 0	0, 0	1, 1	0, 0
MED	SEN	AN	WR	PAL	AMP	0, 0	0, 0	1, 0	0, 0
MED	SEN	AN	WR	PAL	LAT	0, 0	1, 0	1, 1	0, 0
MED	SEN	AN	WR	PAL	VEL	0, 0	0, 0	3, 2	2, 2
MED	SEN	AN	WR	RI	AMP	0, 0	0, 0	1, 0	0, 0
MED	SEN	AN	WR	RI	LAT	0, 0	0, 0	3, 2	3, 2
MED	SEN	AN	WR	RI	VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	AN	WR	TH	AMP	1, 1	0, 0	2, 1	0, 0
MED	SEN	AN	WR	TH	LAT	1, 1	0, 0	3, 2	0, 0
MED	SEN	AN	WR	TH	VEL	0, 0	0, 0	1, 1	1, 1
MED	SEN	OR			AMP	0, 0	1, 0	0, 0	0, 0
MED	SEN	OR			LAT	0, 0	1, 0	0, 0	0, 0
MED	SEN	OR		WR	AMP	0, 0	0, 0	2, 2	0, 0
MED	SEN	OR		WR	LAT	0, 0	0, 0	1, 1	2, 2
MED	SEN	OR		WR	VEL	0, 0	0, 0	2, 2	0, 0
MED	SEN	OR	IN		AMP	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	IN		LAT	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	IN		OTH	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	IN	DAI	VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	IN	PAL	VEL	0, 0	0, 0	0, 0	1, 1

	Nerve	Condu	Patient selection type						
Nerve Tested	Nerve Tested	Configuration	Stimulation	Recording	Parameter	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified
MED	SEN	OR	IN	WR	AMP	4, 3	0, 0	7, 5	2, 2
MED	SEN	OR	IN	WR	LAT	1, 1	0, 0	8, 7	3, 3
MED	SEN	OR	IN	WR	ОТН	2, 2	0, 0	2, 1	1, 1
MED	SEN	OR	IN	WR	PRE	1, 1	0, 0	4, 4	0, 0
MED	SEN	OR	IN	WR	VEL	4, 3	1, 1	8, 7	3, 3
MED	SEN	OR	MI		AMP	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	MI		LAT	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	MI		ОТН	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	MI		VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	MI	MI	AMP	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	MI	MI	VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	MI	PAL	AMP	1, 0	0, 0	0, 0	0, 0
MED	SEN	OR	MI	PAL	VEL	1, 0	0, 0	2, 2	0, 0
MED	SEN	OR	MI	WR	AMP	2, 1	0, 0	3, 3	4, 4
MED	SEN	OR	MI	WR	INCH	1, 1	0, 0	0, 0	2, 2
MED	SEN	OR	MI	WR	LAT	0, 0	0, 0	4, 3	0, 0
MED	SEN	OR	MI	WR	ОТН	1, 1	0, 0	1, 1	0, 0
MED	SEN	OR	MI	WR	PRE	1, 1	0, 0	2, 2	1, 1
MED	SEN	OR	MI	WR	VEL	3, 2	0, 0	5, 5	5, 5
MED	SEN	OR	OTH		VEL	1, 1	0, 0	0, 0	0, 0
MED	SEN	OR	OTH	WR	AMP	0, 0	0, 0	1, 1	0, 0
MED	SEN	OR	OTH	WR	LAT	0, 0	0, 0	2, 2	0, 0
MED	SEN	OR	OTH	WR	VEL	0, 0	0, 0	2, 2	1, 1
MED	SEN	OR	PAL	WR	AMP	0, 0	0, 0	2, 2	1, 1
MED	SEN	OR	PAL	WR	LAT	1, 1	1, 1	11, 11	1, 1
MED	SEN	OR	PAL	WR	OTH	0, 0	0, 0	1, 1	0, 0
MED	SEN	OR	PAL	WR	PRE	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	PAL	WR	VEL	0, 0	0, 0	7, 7	7, 6
MED	SEN	OR	RI		AMP	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	RI		LAT	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	RI		OTH	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	RI		VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	RI	WR	AMP	3, 2	0, 0	3, 2	1, 1
MED	SEN	OR	RI	WR	LAT	1, 1	1, 1	4, 3	1, 1
MED	SEN	OR	RI	WR	OTH	1, 1	0, 0	1, 1	0, 0
MED	SEN	OR	RI	WR	PRE	1, 1	0, 0	1, 1	2, 2
MED	SEN	OR	RI	WR	VEL	2, 1	0, 0	3, 3	2, 2
MED	SEN	OR	TH		AMP	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	TH		LAT	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	TH		OTH	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	TH		VEL	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	TH	ELB	PRE	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	TH	MI	VEL	0, 0	0, 0	0, 0	0, 0
MED	SEN	OR	TH	PAL	VEL	0, 0	0, 0	0, 0	0, 0
MED	SEN	OR	TH	WR	AMP	1, 1	0, 0	3, 3	2, 2
MED	SEN	OR	TH	WR	LAT	0, 0	0, 0	3, 3	0, 0
MED	SEN	OR	TH	WR	OTH	1, 1	0, 0	1, 1	0, 0

	Nerve	Condu	Patient selection type						
Nerve Tested	Nerve Tested	Configuration	Stimulation	Recording	Parameter	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified
MED	SEN	OR	TH	WR	PRE	1, 1	0, 0	1, 1	1, 1
MED	SEN	OR	TH	WR	VEL	1, 1	0, 0	5, 5	2, 2
MED	SEN	OR	WR	ELB	AMP	2, 1	0, 0	0, 0	1, 1
MED	SEN	OR	WR	ELB	OTH	1, 1	0, 0	0, 0	0, 0
MED	SEN	OR	WR	ELB	PRE	0, 0	0, 0	0, 0	1, 1
MED	SEN	OR	WR	ELB	VEL	2, 1	0, 0	0, 0	1, 1
MED	Transcarpal				AMP	0, 0	0, 0	0, 0	1, 1
MED	Transcarpal				LAT	0, 0	0, 0	0, 0	1, 1
RAD	SEN	AN	FOR	TH	LAT	0, 0	0, 0	1, 0	0, 0
RAD	SEN	AN	WR	TH	AMP	1, 1	0, 0	0, 0	0, 0
RAD	SEN	AN	WR	TH	LAT	1, 1	0, 0	2, 2	2, 0
RAD	SEN	AN	WR	TH	VEL	0, 0	0, 0	0, 0	1, 1
RAD	SEN	OR	TH	WR	AMP	1, 0	0, 0	1, 1	0, 0
RAD	SEN	OR	TH	WR	LAT	0, 0	0, 0	1, 1	0, 0
RAD	SEN	OR	TH	WR	PRE	0, 0	0, 0	1, 1	0, 0
RAD	SEN	OR	TH	WR	VEL	1, 0	0, 0	2, 2	1, 1
ULN	MOT				LAT	0, 0	0, 0	0, 0	1, 1
ULN	MOT				OTH	1, 1	0, 0	1, 0	0, 0
ULN	MOT		ELB	ADM	LAT	1, 1	0, 0	0, 0	0, 0
ULN	MOT		ELB	ADM	OTH	1, 1	0, 0	0, 0	0, 0
ULN	MOT		ELB	OTH	AMP	0, 0	0, 0	0, 0	1, 1
ULN	MOT		ELB	OTH	PRE	0, 0	0, 0	0, 0	1, 1
ULN	MOT		ELB	OTH	VEL	0, 0	0, 0	0, 0	1, 1
ULN	MOT		ELB	WR	VEL	1, 1	0, 0	1, 1	1, 1
ULN	MOT		WR		LAT	1, 1	0, 0	1, 1	0, 0
ULN	MOT		WR	1536	VEL	1, 1	0, 0	0, 0	0, 0
ULN	MOT		WR	ADM	AMP	0, 0	0, 0	1, 0	2, 1
ULN	MOT		WR	ADM	LAT	2, 2	1, 1	4, 2	5, 4
ULN	MOT		WR	ADM	OTH	1, 1	0, 0	0, 0	0, 0
ULN	MOT		WR	ADM	VEL LAT	0, 0	0, 0	1, 0	0, 0
ULN	MOT		WR	APB		1, 1	0, 0	0, 0	0, 0
ULN	MOT MOT		WR WR	OTH OTH	AMP LAT	0, 0	0, 0	1, 1	0, 0
ULN	MOT	+	WR	OTH	PRE	0, 0	0, 0	1, 1	0, 0
ULN	MOT	+	WR	PAL	AMP	0, 0	0, 0	1, 1	0, 0
ULN	MOT	+	WR	PAL	LAT	0, 0	0, 0	1, 1	0, 0
ULN	SEN	+	*****	IAL	OTH	0, 0	0, 0	1, 1	0, 0
ULN	SEN	+	WR		AMP	1, 1	0, 0	0, 0	0, 0
ULN	SEN	+	WR	1	LAT	1, 1	0, 0	0, 0	0, 0
ULN	SEN	AN	FOR	LI	LAT	0, 0	0, 0	1, 0	0, 0
ULN	SEN	AN	FOR	RI	LAT	0, 0	0, 0	1, 0	0, 0
ULN	SEN	AN	PAL	LI	LAT	0, 0	0, 0	1, 1	0, 0
ULN	SEN	AN	WR	LI	AMP	0, 0	0, 0	2, 2	1, 1
ULN	SEN	AN	WR	LI	LAT	0, 0	0, 0	2, 2	1, 1
ULN	SEN	AN	WR	LI	VEL	0, 0	0, 0	3, 3	0, 0
ULN	SEN	AN	WR	PAL	LAT	0, 0	0, 0	1, 1	0, 0
ULN	SEN	AN	WR	RI	LAT	0, 0	0, 0	2, 2	4, 2

	Nerve	Condu	Patient selection type						
Nerve Tested	Nerve Tested	Configuration	Stimulation	Recording	Parameter	Complex objective standard	Simple nerve conduction	Symptoms/ presented	Unspecified diagnosis
ULN	SEN	AN	WR	RI	VEL	0, 0	0, 0	0, 0	1, 1
ULN	SEN	OR	LI	WR	AMP	2, 1	0, 0	4, 3	3, 3
ULN	SEN	OR	LI	WR	LAT	1, 1	0, 0	3, 2	1, 1
ULN	SEN	OR	LI	WR	OTH	1, 1	0, 0	1, 0	0, 0
ULN	SEN	OR	LI	WR	PRE	0, 0	0, 0	1, 1	0, 0
ULN	SEN	OR	LI	WR	VEL	2, 1	0, 0	3, 2	3, 3
ULN	SEN	OR	OTH		VEL	1, 1	0, 0	0, 0	0, 0
ULN	SEN	OR	OTH	WR	VEL	0, 0	0, 0	0, 0	1, 1
ULN	SEN	OR	PAL	WR	AMP	0, 0	0, 0	1, 1	0, 0
ULN	SEN	OR	PAL	WR	LAT	0, 0	1, 1	6, 6	0, 0
ULN	SEN	OR	PAL	WR	VEL	0, 0	0, 0	2, 2	1, 1
ULN	SEN	OR	RI	WR	AMP	2, 1	0, 0	2, 1	2, 2
ULN	SEN	OR	RI	WR	LAT	1, 1	1, 1	3, 2	1, 1
ULN	SEN	OR	RI	WR	PRE	0, 0	0, 0	1, 1	1, 1
ULN	SEN	OR	RI	WR	VEL	2, 1	0, 0	2, 2	3, 3
ULN	SEN	OR	WR	ELB	AMP	1, 1	0, 0	0, 0	0, 0
ULN	SEN	OR	WR	ELB	OTH	1, 1	0, 0	0, 0	0, 0
ULN	SEN	OR	WR	ELB	VEL	1, 1	0, 0	0, 0	0, 0

Table 20. Numbers of Studies Reporting Composite Nerve Conduction Tests Across Patient Selection Categories

Legend:

Blank cells—Not reported or not applicable

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

	C	Composite t	est type		Patient selection group				
Nerve for test	Nerve for test	Motor or	Unit of nerve test	Type composite	Complex objective	Simple nerve	Symptoms/ presented	Unspecified diagnosis	
1	2	sensory		7.00	standard	conduction		1.0	
Median	Median	Motor	Amplitude	Difference	0, 0	0, 0	0, 0	1, 0	
Median	Median	Motor	Amplitude	Ratio	1, 1	0, 0	1, 0	0, 0	
Median	Median	Motor	Latency	Difference	0, 0	0, 0	2, 2	2, 2	
Median	Median	Motor	Latency	Ratio	0, 0	0, 0	0, 0	1, 1	
Median	Median	Motor	Velocity	Difference	1, 0	0, 0	0, 0	0, 0	
Median	Median	Sensory	Amplitude	Difference	1, 1	0, 0	2, 2	0, 0	
Median	Median	Sensory	Amplitude	Ratio	1, 1	0, 0	1, 0	1, 1	
Median	Median	Sensory	Latency	Difference	1, 1	0, 0	6, 5	1, 1	
Median	Median	Sensory	Latency	Ratio	0, 0	0, 0	0, 0	1, 1	
Median	Median	Sensory	Velocity	Difference	0, 0	0, 0	2, 2	1, 1	
Median	Median	Sensory	Velocity	Ratio	0, 0	0, 0	4, 4	2, 2	
Median	Radial	Sensory	Latency	Difference	1, 1	0, 0	3, 3	2, 0	
Median	Radial	Sensory	Velocity	Difference	0, 0	0, 0	0, 0	1, 1	
Median	Radial	Sensory	Velocity	Ratio	0, 0	0, 0	1, 1	0, 0	
Median	Ulnar	Motor	Latency	Difference	1, 1	2, 2	3, 3	5, 4	
Median	Ulnar	Motor	Other	Difference	1, 1	0, 0	0, 0	0, 0	
Median	Ulnar	Sensory	Amplitude	Ratio	0, 0	0, 0	2, 2	1, 1	
Median	Ulnar	Sensory	Latency	Difference	1, 1	1, 1	10, 9	5, 3	
Median	Ulnar	Sensory	Velocity	Difference	0, 0	0, 0	1, 1	1, 1	
Median	Ulnar	Sensory	Velocity	Ratio	0, 0	0, 0	1, 1	0, 0	
Radial	Median	Sensory	Velocity	Ratio	0, 0	0, 0	1, 1	0, 0	
Radial	Radial	Sensory	Latency	Difference	0, 0	0, 0	1, 0	0, 0	
Ulnar	Median	Sensory	Velocity	Difference	1, 0	0, 0	0, 0	0, 0	
Ulnar	Median	Sensory	Velocity	Ratio	0, 0	0, 0	1, 1	0, 0	
		•	j	Other Difference	3, 1	0, 0	3, 3	1, 1	
				Other Ratio	0, 0	0, 0	3, 2	1, 1	
				Other Composite	5, 4	0, 0	9, 8	4, 2	

Table 21. Numbers of Articles Reporting Imaging Tests in Patient Selection Categories

Legend:

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Imaging test	Complex objective standard	Simple nerve conduction	Symptoms/presented	Unspecified diagnosis
СТ	0, 0	0, 0	0, 0	2, 0
MRI	2, 0	2, 0	1, 1	5, 2
Ultrasound	1, 0	1, 0	1, 0	3, 3

Summary ROC Meta-Analysis of Diagnostic Test Results

Ideally, a meta-analysis of a test includes only studies that use the same definition of what is to be diagnosed. However, the absence of a gold standard for defining carpal tunnel syndrome resulted in there being as many different definitions of the condition (and therefore of positive cases) as there were studies. Therefore, we could only combine study results by permitting different authors to use different definitions of CTS. Testing for heterogeneity of results helps reduce, but does not eliminate the possibility that different definitions affected study results.

Distal Motor Latency: Patients with Unspecified Diagnosis of CTS v. Normal Controls

While there were 21 studies of distal motor latency (DML) in patient groups coded as "Unspecified diagnosis" that reported some 2 x 2 tables, only five of those studies ultimately could be included in a meta-analysis. Reasons for the exclusion of the others are shown in Table 22. Seven studies did not report any sensitivity or specificity results for the DML measurements, even though they reported them for other tests. Four studies reported sensitivity but not specificity, while one reported specificity but not sensitivity. These studies were excluded because data from both groups are necessary to ensure the validity of the results and because the summary ROC method requires both sensitivity and specificity for each study. The study by Bronson et al. 163 was excluded because DML results were reported for only some of the patients. So et al. ¹⁷³ combined direct measurement of DML with abnormalities in the difference between median and ulnar latency when reporting their results, and we could not isolate results for DML. Charles et al. 170 was excluded because authors reported use of a mean + 2 SD threshold for defining abnormal latency, but the actual threshold reported (4.5 msec) did not agree with their reported results for their control subjects (mean + 2 SD = 4.0 msec). Since the number of controls with latency = 4.5 msec was not reported, we could not derive an internallyconsistent 2 x 2 table from the article, and had to exclude it from analysis. Resende et al. 174 reported patient-level data, but did not report a threshold fordistinguishing normal

Table 22. Distal Motor Latency Studies Excluded from Meta-Analysis

Study	Reason for Exclusion
Pease, 1990 ¹⁷⁷	Did not report sensitivity and specificity for distal motor latency test
Seror, 1998 159	Did not report sensitivity and specificity for distal motor latency test
Rossi, 1994 ¹⁷⁸	Did not report sensitivity and specificity for distal motor latency test
Seror, 1995 ¹⁷⁹	Did not report sensitivity and specificity for distal motor latency test
Lang, 1995 109	Did not report sensitivity and specificity for distal motor latency test
Tzeng, 1990 180	Did not report sensitivity and specificity for distal motor latency test
Mondelli, 2001 ¹⁸¹	Did not report sensitivity and specificity for distal motor latency test
Simovic, 1997 ¹⁸²	Did not report distal motor latency results for control subjects
Simovic, 1999 ¹⁸³	Did not report distal motor latency results for control subjects
Resende, 2000 ¹⁸⁴	Did not report distal motor latency results for control subjects
Lauritzen, 1991 ¹⁸⁵	Did not report distal motor latency results for control subjects
Loscher, 2000 175	Did not report distal motor latency results for CTS patients
Bronson, 1997 163	Selective reporting of distal motor latency results
So, 1989 ¹⁷³	Reported combination test of distal motor latency and other nerve conduction measurements
Charles, 1990 170	Discrepancy in reported threshold
Resende, 2000 174	No diagnostic threshold reported

Meta-analysis of Distal Motor Latency Results in Trials With Table 23. Non-specific Diagnosis of Carpal Tunnel Syndrome Groups

Study	TP	FN	FP	TN	Sen. 95% CI	Spec. 95% CI	PPV 95% CI	NPV 95% CI	Prev.
^a Kimura ¹⁴⁰	105	67	3	119	61.0% 53.4% 68.2%	97.5% 92.9% 99.2%	97.2% 92.0% 99.1%	64.0% 56.7% 70.7%	58.5%
Marin 139	9	5	0	12	64.3% 38.3% 83.9%	100% 75.0% 100%	100% 69.2% 100%	70.6% 46.4% 86.9%	53.8%
Loong ¹⁴¹	17	10	0	30	63.0% 43.9% 78.7%	100% 88.2% 100%	100% 81.0% 100%	75.0% 59.5% 86.0%	47.4%
Plaja ¹⁴²	16	7	0	20	69.6% 48.7% 84.6%	100% 83.3% 100%	100% 80.0% 100%	74.1% 54.9% 87.0%	53.5%
bRosén ¹³⁸	12	29	0	50	29.3% 17.4% 44.8%	100% 92.6% 100%	100% 75.0% 100%	63.3% 52.0% 73.3%	45.1%
Meta-analysis results (mean threshold)					57.1% 49.1% 64.8%	97.9% 97.1% 98.5%			

Sen.-sensitivity, Spec-specificity, PPV-positive predictive value, NPV-negative predictive value, Prev.-prevalence of CTS Confidence intervals on sensitivity, specificity, PPV, NPV calculated by Wilson method⁹⁶

 $[\]underline{\text{Key}}\text{:}\\ \overline{\text{TP-true positive, FN-false negative, FP-false positive, TN-true negative}$

^aCounts for control group (false positive, true negative) estimated by ECRI from threshold reported by authors (mean + 2 SD)

bResults calculated by ECRI from published histogram

The results of this meta-analysis are very similar to the results for the meta-analysis of DML with patient groups with unspecified diagnosis of CTS. The results of both meta-analyses suggest that this test has very high specificity, but only moderate sensitivity.

Table 24. Distal Motor Latency Articles Excluded From Meta-Analysis

Study	Reason for Exclusion
Jackson, 1989 150	Did not report sensitivity and specificity for distal motor latency test
Sener, 2000 ¹⁸⁶	Did not report sensitivity and specificity for distal motor latency test
Schwartz, 1979 ¹⁸⁷	Did not report sensitivity and specificity for distal motor latency test
Escobar, 1985 ¹⁵¹	Did not report sensitivity and specificity for distal motor latency test
Preston, 1992 188	Did not report distal motor latency results for control subjects
Kimura, 1985 ¹⁸⁹	Did not report distal motor latency results for control subjects
Cherniak, 1996 ¹⁹⁰	Used distal motor latency for patient selection
Sheean, 1995 191	Used distal motor latency for patient selection
Foresti, 1996 ¹⁹²	Discrepancies in reported results

Table 25. Meta-analysis of Distal Motor Latency Results in Trials With Patients Presenting with CTS Symptoms

Study	TP	FN	FP	TN	Sen.	Spec.	PPV	NPV	Prev.
					95% CI	95% CI	95% CI	95% CI	
a, b Chang 145	17	26	0	40	39.5%	100%	100%	60.6%	51.8%
					26.1% 54.7%	90.9% 100%	81.0% 100%	48.3% 71.7%	
Kuntzer ¹⁴⁴	47	53	1	69	47.0%	98.6%	97.9%	56.6%	58.8%
					37.3% 56.9%	92.1% 99.8%	88.8% 99.6%	47.5% 65.2%	
^a Murthy ¹⁴³	38	19	2	72	66.7%	97.3%	95.0%	79.1%	43.5%
					53.5% 77.7%	90.5% 99.3%	83.2% 98.6%	69.5% 86.3%	
Cioni ¹⁴⁶	300	75	0	56	80.0%	100%	100%	42.7%	87.0%
					75.6% 83.8%	93.3% 100%	98.7% 100%	34.4% 51.5%	
bMessina ¹²⁰	34	6	1	39	85.0%	97.5%	97.1%	86.7%	50.0%
					70.6% 93.0%	86.8% 99.6%	85.1% 99.5%	73.5% 93.8%	
Melvin ¹⁴⁷	13	4	0	24	76.5%	100%	100%	85.7%	41.5%
					52.2% 90.6%	85.7% 100%	76.5% 100%	68.1% 94.4%	
Loong 148	13	9	0	60	59.1%	100%	100%	87.0%	26.8%
					38.4% 77.0%	93.8% 100%	76.5% 100%	76.8% 93.1%	
^c Atroshi ¹³⁶	25	18	8	52	58.1%	86.7%	75.8%	74.3%	41.7%
					43.0% 71.9%	75.6% 93.2%	58.6% 87.3%	62.7% 83.2%	
Meta-analysis results (mean threshold)					66.0%	98.3%			
•				ĺ	55.7% 75.0%	97.4% 98.9%			

Key:

TP-true positive, FN-false negative, FP-false positive, TN-true negative

Sen.-sensitivity, Spec-specificity, PPV-positive predictive value, NPV-negative predictive value, Prev.-prevalence of CTS

Confidence intervals on sensitivity, specificity, PPV, NPV calculated by Wilson method

^aCounts for control group (false positive, true negative) estimated by ECRI from threshold reported by authors (mean + 2 or 2.5 SD)

bResults calculated by ECRI from published graph

^cOutlier (excluded from meta-analysis results): see text

symptoms of CTS. Although the summary ROC can be extrapolated to a point where sensitivity and specificity are both quite high (i.e., 96%, 96% respectively), in actual practice it is likely that only specificity is so high. Sensitivity was lower than specificity in all five studies.

Table 26. Palmar Sensory Latency Articles Excluded from Meta-analysis

Study	Reason for Exclusion
Gerr, 1998 ³¹	Did not report sensitivity and specificity for palmar sensory latency test
Foresti, 1996 192	Did not report sensitivity and specificity for palmar sensory latency test
Eisen, 1993 ¹⁹³	Did not report sensitivity and specificity for palmar sensory latency test
Mills, 1985 ¹⁹⁴	Did not report sensitivity and specificity for palmar sensory latency test
Kim, 1983 ¹⁹⁵	Did not report sensitivity and specificity for palmar sensory latency test
Andary, 1996 196	Palmar sensory latency results used as patient selection criterion

Table 27. Meta-analysis of Palmar Sensory Latency Results

Study	TP	FN	FP	TN	Sen.	Spec.	PPV	NPV	Prev.
					95% CI	95% CI	95% CI	95% CI	
a, bChang 145	26	17	0	40	60.5%	100%	100%	70.2%	51.8%
					45.3% 73.9%	90.9% 100%	86.7% 100%	57.1% 80.6%	
^c Jackson ¹⁵⁰	91	40	1	37	69.5%	97.4%	98.9%	48.1%	77.5%
					60.9% 76.8%	86.2% 99.5%	93.9% 99.8%	37.0% 59.3%	
^a Murthy ¹⁴³	55	2	2	72	96.5%	97.3%	96.5%	97.3%	43.5%
					87.8% 99.1%	90.5% 99.3%	87.8% 99.1%	90.5% 99.3%	
^a Escobar ¹⁵¹	32	8	2	102	80.0%	98.1%	94.1%	92.7%	27.8%
					64.9% 89.6%	93.1% 99.5%	80.5% 98.4%	86.1% 96.3%	
^c Girlanda ¹⁴⁹	38	37	1	89	50.7%	98.9%	97.4%	70.6%	45.5%
					39.4% 61.9%	93.8% 99.8%	86.5% 99.6%	62.0% 78.0%	
Meta-analys	is resul	lts (me	an thre	shold)	75.8%	97.7%			
					68.8% 81.6%	96.8% 98.4%			

Key:

Sen.–sensitivity, Spec–specificity, PPV–positive predictive value, NPV–negative predictive value, Prev.–prevalence of CTS Confidence intervals on sensitivity, specificity, PPV, NPV calculated by Wilson method

TP-true positive, FN-false negative, FP-false positive, TN-true negative

^aCounts for control group (false positive, true negative) estimated by ECRI from threshold reported by authors (mean + 2 or 2.5 SD)

bResults calculated by ECRI from published graph

cResults calculated by ECRI from published percentages

This left a total of 10 articles for meta-analysis (Table 29). We found significant heterogeneity among the studies' results (Q = 71.4, p < 0.000001). Six studies selected CTS patients using procedures we categorized as "complex objective standard." Analyzing this subgroup separately did not eliminate the heterogeneity (Q = 59.4, p < 0.000001), nor did excluding the one study¹¹¹ that used the reverse Phalen maneuver. (Q = 70.8, p < 0.000001). There were no obvious outliers to explain the heterogeneity, and grouping studies according to criteria that might affect the validity or generalizability of the results (Table 30) did not reduce heterogeneity to statistically non-significant levels. Thus we could not confidently report a single point as the most likely sensitivity and specificity of the test.

The variability of results is shown in Figure 12; sensitivity/specificity covered a large range. We can only conclude that Phalen's maneuver has some ability to distinguish CTS patients from normal controls; the data are too heterogeneous to estimate sensitivity or specificity.

Table 28. Phalen's Maneuver Articles Excluded from Meta-Analysis

Study	Reason for Exclusion
Koris, 1988 ¹⁹⁸	Did not report specificity of Phalen's maneuver
Brahme, 1997 199	Did not report specificity of Phalen's maneuver
Lang, 1995 109	Did not report specificity of Phalen's maneuver
Glass, 1995 ²⁸	Reported results for only 22 of 159 affected hands
Gerr, 1994 ¹⁹⁷	Duplicate publication

Table 29. Diagnostic Trial Results for Phalen's Maneuver

Study	TP	FN	FP	TN	Sen.	Spec.	PPV	NPV	Prev.
					95% CI	95% CI	95% CI	95% CI	
De Smet 101	57	9	4	77	86.4%	95.1%	93.4%	89.5%	44.9%
					75.8% 92.7%	87.8% 98.1%	84.1% 97.5%	81.1% 94.5%	
Durkan ¹⁵⁵	32	14	8	42	69.6%	84.0%	80.0%	75.0%	47.9%
					54.9% 81.1%	71.2% 91.8%	64.9% 89.6%	62.0% 84.6%	
Gellman ¹⁰⁶	45	18	10	40	71.4%	80.0%	81.8%	69.0%	55.8%
					59.0% 81.3%	66.7% 88.9%	69.4% 89.9%	55.9% 79.6%	
a, b Gerr ³¹	48	67	4	11	41.7%	96.7%	92.3%	64.0%	48.3%
				9	33.0% 51.1%	91.8% 98.8%	81.5% 97.0%	56.7% 70.7%	
^b Ghavanini ¹⁵⁴	34	40	17	41	45.9%	70.7%	66.7%	50.6%	56.1%
					34.9% 57.4%	57.7% 81.0%	52.7% 78.2%	39.7% 61.4%	
González del	17	26	20	18	87.0%	90.0%	89.7%	87.4%	50.0%
Pino 104	4			0	81.5% 91.0%	84.9% 93.5%	84.5% 93.3%	82.0% 91.3%	
^a Szabo ¹⁵²	65	22	5	95	74.7%	95.0%	92.9%	81.2%	46.5%
					64.4% 82.8%	88.7% 97.9%	84.1% 97.0%	73.0% 87.3%	
Tetro ¹⁰² 1	58	37	16	80	61.1%	83.3%	78.4%	68.4%	49.7%
					50.8% 70.4%	74.4% 89.6%	67.5% 86.4%	59.3% 76.2%	
Fertl ¹⁵³	50	23	3	36	68.5%	92.3%	94.3%	61.0%	65.2%
					56.9% 78.2%	79.3% 97.4%	84.4% 98.1%	48.0% 72.6%	
^c Werner ¹¹¹	17	14	0	20	54.8%	100%	100%	58.8%	60.8%
					37.5% 71.1%	83.3% 100%	81.0% 100%	41.9% 73.9%	
Meta-analysis results (mean threshold)			NA	NA					

Key:

Sen.—sensitivity, Spec—specificity, PPV—positive predictive value, NPV—negative predictive value, Prev.—prevalence of CTS

Confidence intervals on sensitivity, specificity, PPV, NPV calculated by Wilson method

NA—Results not valid because of excessive heterogenity in study results

TP-true positive, FN-false negative, FP-false positive, TN-true negative

^aResults calculated by ECRI from published percentages

bErrors in published results corrected by ECRI

cTested reverse Phalen's maneuver



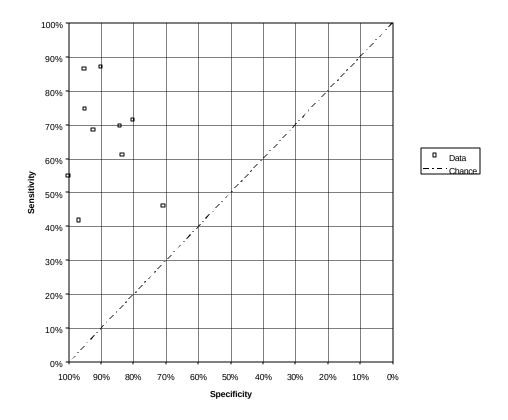


Table 30. Heterogeneity of Diagnostic Trial Results for Phalen's Maneuver

Group	Q (p-value)	
	for larger group	
All articles $(N = 10)$	71.4 (p <0.000001)	
Patients selected with complex objective standard $(N = 6)$ v. other selection	59.4 (p < 0.000001)	
Reverse Phalen's maneuver $(N = 1)$ v. conventional	70.8 (p < 0.000001)	
Not funded by for-profit device or drug manufacturer $(N = 4)$ v. not reported	58.5 (p < 0.000001)	
Reported both inclusion and exclusion criteria $(N = 4)$ v. reported only inclusion criteria	20.5 (p = 0.001)	
Prospective patient selection $(N = 5)$ v. not reported	58.7 (p < 0.000001)	
Comorbidity reported $(N = 1)$ v. not reported	69.9 (p < 0.000001)	
Sex ratios of patients, controls within 20% of each other $(N = 5)$ v. possible sex bias	58.5 (p < 0.000001)	
Mean ages of patients, controls within 5 years $(N = 3)$ v. possible age bias	15.4 (p = 0.017)	
Duration of condition reported $(N = 2)$ v. not reported	48.4 (p < 0.000001)	
Independent reference standard ($N = 4$) v. no independent reference standard reported	48.2 (p < 0.000001)	
Patients given both study test and reference test $(N = 3)$ v. did not do so	49.3 (p < 0.000001)	
Studies done in USA ($N = 6$) v. other countries	58.1 (p < 0.000001)	
Potential selection bias for easy cases $(N = 4)$ v. no bias or not reported	49.3 (p < 0.000001)	

Q—Q-statistic, with probability that variability in study results [D, logit (sensitivity) + logit (specificity)] is the result of random variability within a homogeneous sample of studies.

Tinel's Sign: Combined CTS Groups v. Normal Controls

The evidence base on Tinel's sign comprised 13 studies; three of these reported two CTS groups, for a total of 16 entries in the cross-tabulation. As mentioned in the meta-analysis of Phalen's maneuver, only the later of the duplicate Gerr publications ^{31,197} was included in the analysis, and we pooled patient groups in studies with two CTS groups. Two articles were excluded because they did not report specificity. Exclusions are summarized in Table 31

Eleven studies remained for meta-analysis (Table 32). The meta-analysis found significant heterogeneity among the studies' results (Q = 59.1, p < 0.000001). All but two studies (De Smet et al. ¹⁰¹ and Seror et al. ¹⁵⁶) selected CTS patients using procedures we categorized as "complex objective standard." Excluding those studies from the analysis did not substantially reduce the heterogeneity (Q = 46.7, p < 0.000001).

The heterogeneity is evident in Figure 13. Sensitivity/specificity results are widely dispersed in the graph, and there is no pattern of results that is obvious on inspection. The data suggest that Tinel's sign has some ability to diagnose CTS, but the sensitivity and specificity of the test are uncertain. However, the sensitivity of the test appears to be low.

To see whether other factors, particularly those relating to the validity or generalizability of results, could explain the observed heterogeneity, we repeated the heterogeneity tests for groups defined by reporting criteria in Table 13 and Table 15. The results of those analyses are shown in Table 33. Significant heterogeneity remained regardless of the criteria used to group trials. Therefore none of these criteria are sufficient to explain the heterogeneity that prevents us from meta-analyzing the results.

Table 31. Tinel's Sign Articles Excluded from Meta-analysis

Study	Reason for Exclusion		
Brahme, 1997 ¹⁹⁹	Did not report specificity of Tinel's sign		
Lang, 1995 109	Did not report specificity of Tinel's sign		
Gerr, 1994 ¹⁹⁷	Duplicate publication		

Table 32. Diagnostic Trial Results for Tinel's Sign

Study	TP	FN	FP	TN	Sen.	Spec.	PPV	NPV	Prev.
					95% CI	95% CI	95% CI	95% CI	
De Smet ¹⁰¹	14	17	0	81	45.2%	100%	100%	82.7%	27.7%
					28.9% 62.5%	95.3% 100%	77.8% 100%	73.8% 89.0%	
Durkan ¹⁵⁵	26	20	10	40	56.5%	80.0%	72.2%	66.7%	47.9%
					42.0% 70.0%	66.7% 88.9%	55.7% 84.3%	53.8% 77.5%	
Gellman ¹⁰⁶	29	37	3	47	43.9%	94.0%	90.6%	56.0%	56.9%
					32.4% 56.2%	83.5% 98.0%	75.4% 96.8%	45.1% 66.3%	
Gelmers ²⁹	20	27	11	32	42.6%	74.4%	64.5%	54.2%	52.2%
					29.3% 57.0%	59.4% 85.2%	46.6% 79.1%	41.4% 66.5%	
a, bGerr ³¹	8	50	2	121	13.8%	98.4%	80.0%	70.8%	32.0%
					7.1% 25.2%	94.1% 99.6%	48.4% 94.5%	63.4% 77.2%	
Ghavanini ¹⁵⁴	24	52	9	49	31.6%	84.5%	72.7%	48.5%	56.7%
					22.1% 42.9%	72.8% 91.7%	55.4% 85.1%	38.8% 58.3%	
González del	42	87	6	194	32.6%	97.0%	87.5%	69.0%	39.2%
Pino ¹⁰⁴					24.9% 41.2%	93.5% 98.6%	75.0% 94.2%	63.3% 74.3%	
^a Seror ¹⁵⁶	63	37	18	22	63.0%	55.0%	77.8%	37.3%	71.4%
					53.0% 72.0%	39.5% 69.6%	67.4% 85.6%	25.9% 50.3%	
Stewart 157	23	28	15	37	45.1%	71.2%	60.5%	56.9%	49.5%
					32.0% 58.9%	57.4% 81.8%	44.4% 74.6%	44.6% 68.5%	
^a Szabo ¹⁵²	56	31	1	99	64.4%	99.0%	98.2%	76.2%	46.5%
					53.7% 73.8%	94.4% 99.8%	90.5% 99.7%	68.0% 82.8%	
^a Tetro ¹⁰²	70	25	9	87	73.7%	90.6%	88.6%	77.7%	49.7%
					63.8% 81.6%	82.9% 95.1%	79.5% 94.0%	68.9% 84.5%	
Meta-analys	sis resul	ts (mea	an thres	shold)	NA	NA			

TP-true positive, FN-false negative, FP-false positive, TN-true negative

Confidence intervals on sensitivity, specificity, PPV, NPV calculated by Wilson method

Sen.-sensitivity, Spec-specificity, PPV-positive predictive value, NPV-negative predictive value, Prev.-prevalence of CTS

NA—Results not valid because of excessive heterogenity in study results

^aResults calculated by ECRI from published percentages

bErrors in published results corrected by ECRI

Table 33. Heterogeneity of Diagnostic Trial Results for Tinel's Sign

Group	Q (p-value)
	for larger group
All articles $(N = 11)$	59.1 (p < 0.000001)
Patients selected with complex objective standard $(N = 9)$ v. other selection	46.1 (p < 0.000001)
Not funded by for-profit device or drug manufacturer $(N = 5)$ v. not reported	10.7 (p = 0.057)
Reported both inclusion and exclusion criteria $(N = 6)$ v. reported only inclusion criteria	30.2 (p = 0.000013)
Prospective patient selection $(N = 4)$ v. not reported	16.6 (p = 0.011)
Comorbidity reported $(N = 2)$ v. not reported	51.4 (p < 0.000001)
Mean ages of patients, controls within 5 years $(N = 6)$ v. possible age bias	37.8 (p < 0.000001)
Possible sex bias $(N = 3)$ vs. sex ratios of patients, controls within 20% of each other	52.8 (p < 0.000001)
(N=8)	
Duration of condition reported $(N = 2)$ v. not reported	50.6 (p <0.000001)
Independent reference standard $(N = 6)$ v. no independent reference standard reported	16.5 (p = 0.005545)
Patients given both study test and reference test $(N = 3)$ v. did not do so	51.6 (p < 0.000001)
Studies done in USA $(N = 5)$ v. other countries	22.3 (p = 0.000454)
Potential selection bias for easy cases $(N = 4)$ v. no bias or not reported	41.9 (p < 0.000001)

Q—Q-statistic, with probability that variability in study results [D, logit (sensitivity) + logit (specificity)] is the result of random variability within a homogeneous sample of studies.

Articles on Carpal Tunnel Syndrome Screening

Screening tests are intended to identify persons at risk of developing a condition in the future, not those who already have the condition. Because there is no agreement on what constitutes screening for CTS, we accepted any studies so described by their authors as screening studies. There were 28 articles described by their authors as screening studies. Two (Bland²⁰⁰ and Rosen²⁰¹) were excluded from this analysis because they required all participants to be symptomatic. Two^{202,203} were sequential reports on the same study. Therefore, 25 studies (Table 34) were included in the analysis of screening of carpal tunnel syndrome. Twenty-two of the studies screened workers at risk, and the remaining three studies screened the general population; the table is stratified according to these two categories.

The reported methods of diagnosis in the 28 screening studies appear in Table 35. The most common diagnostic criteria were symptoms (12 studies, 43%) and the difference between median and ulnar sensory tests (9 studies, 32%). Thirteen studies (46%) used both clinical criteria and nerve conduction criteria, three studies (11%) used nerve conduction criteria only, and no studies used clinical criteria only. The table demonstrates the variability in authors' methods for screening for CTS. As with the diagnostic articles on CTS, we tabulated the number of screening articles reporting use of each particular test (Table 36, Table 37, Table 38, Table 39, Table 40). In no case were there sufficient articles reporting a particular test to meet our a priori criteria for meta-analyzing their data.

The presence of symptoms and the presence of a positive nerve conduction test appeared to be independent of each other in the screening studies. Figure 14 plots the prevalence of symptoms on the horizontal axis and the prevalence of positive nerve conduction tests on the vertical axis. We could only plot the 15 studies that reported both variables. The

"maximum latency difference" test, which is a variation of the inching test. We reanalyzed this data: the resulting sensitivities and specificities at different threshold values are shown in Table 43 and an ROC curve fitted to the data using the logit regression method is shown in Figure 15. While it is clear that this test had a significant ability to predict future CTS in this screening population, this is just one of several nerve conduction tests done in this study, and the possibility of a chance result cannot be discounted. Independent confirmation of this finding would be necessary for us to conclude that this is an effective predictive test. Reanalysis of the unpublished results from this study could verify whether or not other nerve conduction tests also predict future CTS, and could help clinicians decide which test is most effective.

Table 34. Articles Described as Screening Studies

Article	N Population		Symptoms	Positive NCS	Symptoms & Positive NCS
	Wo	orkers-at-risk screening studies for	r carpal tunnel	syndrome	•
Kearns, 2000	45	Pork processors	NR	NR	NR
Missere, 1999 205	45	Meat manufacturers	NR	^a 28.9%	NR
Nathan, 1998 202	283	Steel mill workers, food processors, electronics workers, and plastics workers	12.9%	43.0%	8.2%
Tan, 1998 ²⁰⁶	64	Carpet weavers	NR	NR	NR
Werner, 1998 207	119	Automobile parts manufacturers	NR	27%	^b 20.2%
	98 77 64 164	Furniture manufacturers Paper containers manufacturers Automobile parts manufacturers Clerical insurance workers	NR NR NR NR NR	26% 34% 30% 15% 28%	b 10.2% b 14.3% b 17.2% b 11.0% b 9.4%
Franzblau, 1997 ²⁰⁸	202 Spark plugs manufacturers 148 Automobile parts manufacturers		41%	NR	9.4% NR
Jeng, 1997 ²⁰⁹	27	Food processors	48.8%	34.1%	22.0%
Werner, 1997	59	Manufacturing workers and clerical workers	11.1%	45.4%	5.6%
Bingham, 1996 ²¹¹	am, 102 Applicants for jobs in meat		^c 6.0%	^a 17.4%	^c 1.8%
Murata, 1996 164	27	Data entry operators	NR	37%	NR
Pierre-Jerome, 1996 ²¹²	24	Floor cleaners	NR	NR	NR
Werner, 1995	167	Automobile parts manufacturers	19.8%	24.6%	9.0%
Young, 1995	157	Poultry processors	70%b	31%	NR
Franzblau, 1994 ¹¹³	84	Automobile parts manufacturers	21.4%	19.3%	8.40%
Kirschberg, 112 Poultry processors			22.3%	29.5%	17.0%

Article	N	Population	Symptoms	Positive NCS	Symptoms & Positive NCS
	Wo	rkers-at-risk screening studies for	r carpal tunnel	syndrome	
Nathan, 1994 101		Japanese furniture factory workers	^{a, b} 4.5%	ь 17.8%	^b 2.0%
	316	Steel mill workers, food processors, electronics workers, and plastics workers	^{a, b} 23.4%	^b 22.0%	^b 8.3%
Nilsson, 1994 216	61	Office workers	NR	33%	NR
	58	Truck assemblers	NR	40%	NR
	56	Platers	NR	55%	NR
Werner, 1994 217	130	Automobile parts manufacturers	27.7%	^d 20.2%	NR
Johnson, 1993 167	1993 184 Poultry processors		^{a, b} 37.3%	^{a, b} 19.2%	^{a, b} 6.0%
processors, electronics wo plastics workers, aluminur reduction workers, and cal		Steel mill workers, meat/food processors, electronics workers, plastics workers, aluminum reduction workers, and cable plant workers.	^{a, b} 51.0%	^{a, b} 33.6%	^{a, b} 19.8%
Grant, 1992 219	63	Manufacturing plant workers	^a 25.4%	NR	NR
Jetzer, 1991 168	39	Computer assemblers	NR	NR	NR
	100	Meat processors	NR	NR	NR
	284	Keyboard operators	NR	NR	NR
General populati	on scre	eening studies for carpal tunnel sync	lrome		•
Atroshi, 1999 246 General population 6		14.4%	° 22.3%	^c 6.6%	
Ferry, 1998 ²²¹	648	General population	18.5%	17.4%	7.7%
DeKrom, 1990	500	General population	13.8%	NR	^c 7.8%

NR-Not reported
NCS-Nerve conduction studies

^aBased on hands instead of participants
^bCalculated by ECRI based on information reported in the article
^cEstimated by ECRI based on information reported in the article
^dPrevalence of positive NCS in the study by Werner²¹⁷ was based on 129 participants.

 Table 35. Definitions of CTS Reported in Screening Articles

Author, Clinical findings				Nerv	e cond	duction	n studie	s	Comments	
Year	SYM CLN OTH		DML				MOT	OTH		
			CLN				DIF	DIF	NCS	
Bland, 2000 200	?	?	?	V	?	?	?	?	V	If tests equivocal, authors measured sensory potential or inching test
Kearns, 2000 204	?	?	?	?	?	?	?	?	?	NR
Atroshi, 1999 220	V	V	?	?	?	?	V	?	?	
Missere, 1999 205	?	?	?	?	?	?	?	?	V	
Ferry, 1998	?	?	?	?	?	?	?	?	?	NR
Nathan, 1998 202		?	?	?	V	V	?	?	V	
Rosen, 1998	?	?	?	?	?	?	?	?	?	NR
Tan, 1998 ²⁰⁶	?	?	?	?	?	?	?	?	?	NR
Werner, 1998 207		?	?	?	?	?	V	?	?	
Franzblau, 1997 ²⁰⁸	?	?	?	?	?	?	?	?	?	NR
Jeng, 1997 ²⁰⁹	\checkmark	?	?	$\overline{\checkmark}$?	V	?	?	
Werner, 1997		?	?	?	?	?	V	?	?	
Bingham, 1996 ²¹¹	?	?	?	?	?	?	?	?	?	NR
Murata, 1996	?	?	?	?	?	?	?	?	?	NR
Pierre- Jerome, 1996	?	?	?	?	?	?	?	?	?	NR
Werner, 1995	V	?	?	?	?	?	V	?	?	
Young, 1995	?	?	?	?	?	?	?	?	?	NR
Franzblau, 1994 ¹¹³	V	?	?	?	?	?	V	?	?	
Kirschberg, 1994 ²¹⁴	V	V	V	V	?	V	V	?	V	
Nathan, 1994 215	7	?	?	?	V	V	?	?	V	
Nilsson, 1994 ²¹⁶	?	?	?	?	?	?	?	?	?	NR
Werner, 1994 217	?	?	?	?	?	?	?	?	?	NR
Johnson, 1993 167	?	?	?	?	?	?	?	?	?	NR
Nathan, 1993 218	\square	?	?	?	V	V	?	?	$\overline{\checkmark}$	
Grant, 1992 ²¹⁹	?	?	?		\checkmark	?	$\overline{\checkmark}$	$\overline{\checkmark}$?	

Author,	Clinical findings			Nerve conduction studies				Comments		
Year	SYM	CLN	OTH	DML	DSL	PAL	SEN	MOT	OTH	
			CLN				DIF	DIF	NCS	
Jetzer, 1991 168	V	?	?	?	?	?	?	?	?	Or positive NCS (tests not reported)
DeKrom, 1990 ²²²	V	?	?	V	?	?	$\overline{\mathbf{V}}$?	?	
Welch, 1973	?	?	?	?	?	?	?	?	?	NR
Totals	12	2	1	5	5	4	9	1	6	

Kev

SYM—Were positive symptoms included in the author's method of diagnosis?

CLN—Was a positive clinical exam included in the author's method of diagnosis?

OTH CLN —Were other clinical findings included in the author's method of diagnosis?

DML—Was distal motor latency included in the author's method of diagnosis?

DSL—Was distal sensory latency included in the author's method of diagnosis?

PAL—Was palmar sensory latency included in the author's method of diagnosis?

SEN DIF—Was the difference between median and ulnar sensory studies included in the author's method of diagnosis?

MOT DIF—Was the difference between median and ulnar motor studies included in the author's method of diagnosis?

OTH NCS—Were other nerve conduction studies included in the author's method of diagnosis?

NR—Method of diagnosis was not reported

Figure 14. Association of Symptoms with Positive NCS Findings in Screening Studies

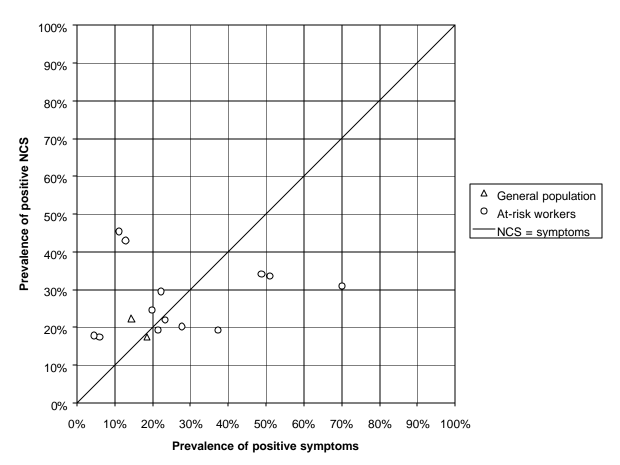


Table 36. Signs and Symptoms Reported in Screening Articles

Legend:

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Sign/symptom	Number of articles reporting
Clinical exam and history	1, 0
Durkan compression	1, 1
Flick sign	1, 1
Flick: Does shaking alleviate night symptoms?	1, 1
Gilliat tourniquet	1, 1
Grip strength	2, 0
Hypalgesia	1, 0
Hyperpathia	1, 0
Lateral pinch strength	1, 0
Luthy's test	1, 1
Night symptoms	1, 1
Opponens pollicus weakness	1, 1
Phalen's/reverse Phalen's	3, 2
Right or left hand worse? Or bilateral?	1, 1
Signs	1, 0
Symptoms measured systematically	15, 7
Symptoms	2, 0
Symptoms and signs	1, 0
Thenar atrophy	1, 1
Thenar weakness	1, 1
Three-point pinch strength	1, 0
Tinel's	3, 2
When are symptoms worse?	1, 1
Which fingers are worst affected?	1, 1

 Table 37. Sensory Tests Reported in Screening Articles

Legend:

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Sensory test	Number of articles reporting
Current perception	1, 1
Gap detection test	1, 1
Semmes-Weinstein monofilament	1, 0
Tactile discrimination	1, 1
Vibrometer	6, 3

Table 38. Nerve Conduction Tests Reported in Screening Articles

Legend:

Nerve tested: MED-median, RAD-radial, ULN-ulnar

Nerve tested: MOT-motor, SEN-Sensory

Configuration (not applicable to motor nerve tests: OR-orthodromic, AN-antidromic

Stimulation/measurement sites: ELB-elbow, FOR-forearm, WR-wrist, PAL-palm, IN-index finger, MI-middle finger, RI-ring finger, LI-little finger, APB-abductor policis brevis, ADM-abductor digiti minimi, OTH-other

Measured parameter: LAT-latency, AMP-amplitude, VEL-velocity, INCH-inching, OTH-other

Blank cells—characteristic not reported First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Numeric entries—Total number of articles, articles from which sensitivity and specificity can be calculated

Nerve tested		Configuration	Stimulation	Measurement	Parameter	Number of articles	
			site	site	measured	reporting	
MED	MOT				LAT	2, 0	
MED	MOT		FOR	APB	LAT	1, 1	
MED	MOT		WR	APB	LAT	4, 2	
MED	MOT		WR	APB	VEL	1, 1	
MED	MOT		WR	OTH	AMP	1, 0	
MED	MOT		WR	OTH	LAT	3, 2	
MED	MOT		WR	OTH	VEL	1, 0	
MED	SEN				AMP	1, 0	
MED	SEN				LAT	4, 0	
MED	SEN				OTH	1, 1	
MED	SEN	AN			LAT	1, 1	
MED	SEN	AN	PAL	IN	VEL	1, 1	
MED	SEN	AN	PAL	MI	AMP	1, 1	
MED	SEN	AN	PAL	MI	VEL	1, 1	
MED	SEN	AN	WR	IN	AMP	2, 2	
MED	SEN	AN	WR	IN	LAT	5, 3	
MED	SEN	AN	WR	IN	VEL	1, 1	
MED	SEN	AN	WR	MI	AMP	1, 1	
MED	SEN	AN	WR	MI	INCH	3, 1	
MED	SEN	AN	WR	MI	VEL	1, 1	
MED	SEN	AN	WR	OTH	LAT	3, 1	
MED	SEN	AN	WR	PAL	VEL	2, 2	
MED	SEN	AN	WR	RI	LAT	1, 1	
MED	SEN	OR	IN	WR	LAT	1, 1	
MED	SEN	OR	IN	WR	VEL	1, 0	
MED	SEN	OR	PAL	WR	LAT	5, 2	
MED	SEN	OR	WR	ELB	VEL	1, 1	
ULN	MOT				LAT	1,0	
ULN	MOT		WR	ADM	LAT	1, 0	
ULN	SEN				LAT	2, 0	
ULN	SEN	AN			LAT	1, 1	
ULN	SEN	AN	WR	LI	AMP	2, 2	
ULN	SEN	AN	WR	LI	LAT	4, 2	
ULN	SEN	AN	WR	RI	LAT	1, 1	
ULN	SEN	OR	Ц	WR	LAT	1, 0	
ULN	SEN	OR	LI	WR	VEL	1,0	
ULN	SEN	OR	PAL	WR	LAT	3, 2	

119

Table 39. Composite Nerve Conduction Tests Reported in Screening Articles

Legend:

Nerves: MED—median, ULN—Ulnar

Measured parameter: LAT-latency, VEL-velocity

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

First	Second	Motor or	Parameter	Combination	Number of articles
nerve	nerve	Sensory	Measured		reporting
MED	MED	SEN	VEL	Ratio	1, 1
MED	ULN	MOT	LAT	Difference	2, 0
MED	ULN	SEN	LAT	Difference	11, 6
ULN	MED	SEN	LAT	Difference	1, 0
				Other composite	7, 3

Table 40. Imaging Tests Reported in Screening Articles

Legend:

First entry in cell—Total number of articles

Second entry in cell—Number of articles with derivable sensitivity and specificity

Imaging modality	Number of articles reporting
CT	1, 0
MRI	1, 0
Ultrasound	1, 1

Table 41. Definitions of CTS Reported in Screening Articles

Article	Method of diagnosis used to determine patient condition
Bland, 2000 ²⁰⁰	Median and ulnar sensory conduction (velocity?), DML to APB. Sensory potential or segmental study of conduction used if previous tests equivocal. Threshold 2.5 SD from the mean.
Kearns, 2000	Not reported
Atroshi, 1999 220	Two definitions: 1) Symptoms and positive clinical exam. Symptoms were pain, numbness and/or tingling in 2 or more of the first 4 fingers at least twice weekly during the preceding 4 weeks, as stated on a questionnaire. Clinical exam required the presence of nocturnal and/or activity-related numbness and/or tingling involving the palmar aspects of at least 2 of the first 4 fingers. The presence of median nerve sensory and/or motor deficit was supportive of the diagnosis but not necessary. 2) Symptoms and positive clinical exam and positive nerve conduction. Included the same definitions as above, and in addition required a difference of 0.8 ms or more between the median sensory latency (middle finger to wrist) and the ulnar sensory latency (little finger to wrist).
Missere, 1999 205	SCV <42.5 m/s as measured by the nerve conduction inching test.
Ferry, 1998 ²²¹	Not reported
Nathan, 1998 202	Symptoms and abnormal nerve conduction. Symptoms defined as positive when the patient has either one of two sets of symptoms: 1) Two or more specific CTS symptoms such as numbness, tingling, nocturnal awakening occurring at least twice per month in the median nerve distribution. 2) One specific CTS symptoms and two or more nonspecific symptoms such as pain, tightness, clumsiness occurring at least
	twice per month in the median nerve distribution. NCS was defined as abnormal when a patient had any of the following three abnormalities: 1) Maximum latency difference = 0.4 ms in the orthodromic inching test. 2) Antidromic wrist-to-digit sensory latency >3.6 ms. 3) Orthodromic palm to wrist sensory latency >2.2 ms
Rosen, 1998	Not reported
Tan, 1998 ²⁰⁶	Not reported
Werner, 1998 207	Nerve conduction abnormality defined as a difference >0.5 ms between median and ulnar antidromic sensory latencies to index and little fingers, respectively. Symptom abnormality defined as numbness, tingling, burning, or pain in the wrist, fingers, or hand.
Franzblau, 1997 ²⁰⁸	Not reported
Jeng, 1997 ²⁰⁹	Two definitions: One required both symptoms and abnormal conduction, and the other required either symptoms or abnormal nerve conduction: Symptoms: tingling, numbness, pain, perceived weakness, and clumsiness. Nerve conduction was abnormal on any of the following three tests: 1) DML >4.5 ms. 2) Antidromic sensory latency from index finger >3.7 ms. 3) Difference between median palm-to-wrist latency and ulnar palm-to-wrist latency >0.5 ms.
Werner, 1997	Difference between median and ulnar sensory latency >0.5 ms, and symptoms.
Bingham, 1996 ²¹¹	Not reported
Murata, 1996	Not reported
Pierre-Jerome, 1996 ²¹²	Not reported
Werner, 1995 213	Symptoms and abnormal NCS. Positive symptoms were defined as any of the following: numbness, tingling, buning, pain, or nocturnal paresthesia in the hand. Abnormal CTS was defined as a difference greater than 0.5 ms between the median and ulnar sensory antidromic latencies.
Young, 1995	Not reported

Article	Method of diagnosis used to determine patient condition
Franzblau, 1994 ¹¹³	Symptoms and abnormal nerve conduction. Positive symptoms was defined as having both 1) numbness, tingling, burning, or pain in the fingers, hand, wrist, or forearm and 2) nocturnal occurrence of above symptoms. Abnormal nerve conduction was defined as a difference >0.5 between median sensory antidromic wrist-to-index latency and ipsilateral ulnar sensory antidromic wrist-to-little-finger latency.
Kirschberg, 1994 ²¹⁴	Clinical CTS: One or more of the following 7 findings: 1) nocturnal paresthesia of the hand, relieved by shaking; 2) sensory symptoms in the specific distribution of the median nerve; 3) specific median nerve sensory loss; 4) positive Phalen's sign; 5) Positive Tinel's sign; 6) Thenar atrophy; 7) Thenar weakness. Electrodiagnostic CTS (using Mayo Clinic criteria) involved any of the following 4 findings: 1) Median DML >4.6 ms; 2) Median palmar sensory latency >2.2 ms; 3) Difference >0.2 ms between median and ulnar palmar latencies; 4) Difference >1.8 ms between median and ulnar latencies.
Nathan, 1994 ²¹⁵	Symptoms and abnormal nerve conduction. Symptoms defined as positive when the patient has either one of two sets of symptoms: 1) Two or more specific CTS symptoms such as numbness, tingling, nocturnal awakening occurring at least twice per month in the median nerve distribution 2) One specific CTS symptom and two or more nonspecific symptoms such as pain, tightness, clumsiness occurring at least twice per month in the median nerve distribution. NCS was defined as abnormal when a patient had any of the following three abnormalities: 1) Maximum latency difference = 0.4 ms in the orthodromic inching test. 2) Antidromic wrist-to-digit sensory latency >3.6 ms 3) Orthodromic palm to wrist sensory latency >2.2 ms
Nilsson, 1994 216	Not reported
Werner, 1994	Not reported
Johnson, 1993 167	Not reported
Nathan, 1993 218	Symptoms and abnormal nerve conduction. Symptoms defined as positive when the patient has either one of two sets of symptoms: 1) Two or more specific CTS symptoms such as numbness, tingling, nocturnal awakening occurring at least twice per month in the median nerve distribution 2) One specific CTS symptoms and two or more nonspecific symptoms such as pain, tightness, clumsiness occurring at least twice per month in the median nerve distributionNCS was defined as abnormal when a patient had any of the following three abnormalities: 1) Maximum latency difference = 0.4 ms in the orthodromic inching test. 2) Antidromic wrist-to-digit sensory latency >3.6 ms 3) Orthodromic palm to wrist sensory latency >2.2 ms
Grant, 1992 ²¹⁹	Median DML >4.5 ms or median DSL >3.5 ms or median-ulnar DML difference >1.2 ms or median-ulnar DSL difference >0.5 ms
Jetzer, 1991 168	Symptoms and either positive EMG or recent prior carpal tunnel surgery.
DeKrom, 1990	Nocturnal paresthesia at least twice a week and either DML >4.5 ms or a difference >0.4 ms between median and ulnar antidromic latencies to the ring finger.
Welch, 1973	Not reported

Table 42. Screening Articles Reporting Longitudinal Results

Article	N	Population	Selection	Followup
Kearns, 2000 ²⁰⁴	45	Porkprocessors	Starting employment	42-83 days, mean 64
Nathan, 1998 ²⁰² 203 218	283	Various manufac- turing and clerical	Randomly-selected workers	11 years
Werner, 1997	NR, though over 700	Various manufac- turing and clerical	NCS positive workers and matched controls	10 to 24 months
Johnson, 1993 167	184	Meat processors	Mostly new employees	Not reported, but few followed more than 3 months

Table 43. Prediction of Future CTS by Maximum Latency Difference

MLD result	Future CTS	No CTS	Threshold	Sensitivity	Specificity	PPV	NPV
<0.28 ms	3	129	0.20	90.9%	29,9%	9.0%	97.7%
0.28-0.35 ms	11	211	0.28 ms	76.1% 96.9%	25.7% 34.5%	6.4% 12.7%	93.4% 99.2%
0.20-0.33 ms	11	211				17.3%	
0.36-0.43 ms	7	56	0.36 ms	57.6% 40.5% 73.0%	78.9% 74.7% 82.5%	11.2% 25.6%	96.0% 93.4% 97.7%
	·					25.5%	
0.44-0.51 ms	5	20	0.44 ms	36.4% 22.0% 53.7%	91.9% 88.8% 94.1%	15.1% 39.8%	95.0% 92.4% 96.7%
0 0.01 1110						31.8%	
>0.51 ms	7	15	0.52 ms	21.2% 10.5% 38.1%	96.5% 94.3% 97.9%	16.1% 53.1%	94.1% 91.5% 96.0%

Data from Nathan et al., 1998 202
Future CTS—Patients developed CTS during the 11-year followup periof No CTS—Patients did not develop CTS during followup period.

Table 44. Carpal Tunnel Syndrome-Study Design

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?
Finsen, 2001 224	$\overline{\square}$			V			Single	1	68	0	0	Prospective	Counts	No: thresholds not reported
Mondelli, 2001 181			V			V	Single	1	20	1	19	Not reported	Counts	Calculated by ECRI
Atroshi, 2000 225			V				Single	1	262	1	125	Prospective	Summary	No: only summary statistics reported
Bland, 2000 200						V	Single	1	8223	1	3533	Retrospective	Counts	Reported by authors
Cuturic, 2000 ²²⁶			V				Single	1	19	1	16	Prospective	Patient level	Calculated by ECRI
Kearns, 2000 ²⁰⁴			V	V			Single	1	45	0	0	Prospective	Summary	No: only summary statistics reported
Loscher, 2000 175			$\overline{\mathbf{A}}$	Ø			Single	2	NR	1	87	Prospective	Counts	Reported by authors
Montagna, 2000 227						V	Single	1	30	1	15	Not reported	Counts	Reported by authors
Nakamichi, 2000 ²²⁸			V		V		Single	1	125	1	200	Not reported	Summary	No: only summary statistics reported
Raudino, 2000 ²²⁹	\square						Single	1	83	0	0	Not reported	Counts	Reported by authors
Resende, 2000 184			Ŋ	V			Single	1	32	1	20	Not reported	Patient level	Calculated by ECRI
Resende, 2000 174			Ŋ				Single	1	20	1	20	Not reported	Patient level	Calculated by ECRI
Sener, 2000 186	V		Ŋ			Ŋ	Single	1	31	1	21	Not reported	Counts	Calculated by ECRI
Seror, 2000 158			Ŋ	V			Single	1	20	1	20	Not reported	Counts	Reported by authors
Stalberg, 2000 ²³⁰				Ø			Single	1	136	1	32	Not reported	Counts	Reported by authors
Weber, 2000 108	Ø	Ø		Ø			Single	1	53	1	26	Not reported	Counts	Reported by authors
Atroshi, 1999 220			V	V			Single	1	2466	0	0	Prospective	Counts	No: only one patient group
Burke, 1999 ²³¹	\square		Ø				Multiple (<5)	1	186	0	0	Prospective	Counts	Calculated by ECRI
Duncan, 1999 ²³²					V		Single	1	68	1	36	Prospective	Counts	Reported by authors
Kabiraj, 1999 ²³³			V	Ø			Single	1	31	1	38	Not reported	Counts	Calculated by ECRI
Lee, 1999 ²³⁴					V		Single	1	50	1	28	Prospective	Counts	Reported by authors
Missere, 1999 ²⁰⁵			V		V		Single	1	45	0	0	Not reported	Counts	Reported by authors
Mongale, 1999 ²³⁵					V		Single	1	8	2	16	Not reported	Summary	No: only summary statistics reported

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?
Murthy, 1999 143			$\overline{\square}$			$\overline{\mathbf{A}}$	Single	1	84	1	37	Not reported	Counts	Reported by authors
Rudolfer, 1999 ²³⁶				V			Single	1	937	0	0	Retrospective	Counts	Calculated by ECRI
Sander, 1999 ²³⁷			\square			$\overline{\mathbf{A}}$	Single	1	59	1	34	Prospective	Counts	Reported by authors
Simovic, 1999 183			Ø	$\overline{\mathbf{A}}$			Single	2	66	1	19	Prospective	Counts	Reported by authors
Szabo, 1999 152	Ø	$\overline{\checkmark}$	Ø			V	Single	1	50	2	100	Prospective	Counts	Reported by authors
Thonnard, 1999 117		$\overline{\checkmark}$	Ø			$\overline{\mathbf{A}}$	Single	1	11	1	10	Prospective	Summary	No: only summary statistics reported
Wang, 1999 ²³⁸			Ø	$\overline{\mathbf{A}}$			Single	1	12	1	12	Prospective	Summary	No: only summary statistics reported
Aurora, 1998 ²³⁹			Ø				Single	1	19	1	20	Not reported	Summary	No: only summary statistics reported
Ferry, 1998 ²²¹	Ø		Ø				Single	1	648	0	0	Prospective	Counts	Reported by authors
Fertl, 1998 153	Ø		Ø			V	Single	1	47	1	20	Prospective	Counts	Reported by authors
Gerr, 1998 31	Ø	$\overline{\checkmark}$	Ø	V		V	Single	1	60	1	59	Not reported	Counts	Reported by authors
Ghavanini, 1998 154	$\overline{\square}$	$\overline{\checkmark}$	$\overline{\square}$				Single	1	74	1	58	Prospective	Counts	Reported by authors
Girlanda, 1998 149	Ø		Ø	$\overline{\mathbf{A}}$			Single	1	41	1	45	Not reported	Counts	Reported by authors
Kabiraj, 1998 ²⁴⁰			Ø	V			Single	1	72	1	65	Retrospective	Summary	No: only summary statistics reported
Kleindienst, 1998 241					N		Single	1	77	1	18	Prospective	Summary	No: only summary statistics reported
Luchetti, 1998 ²⁴²						$\overline{\mathbf{A}}$	Single	1	39	1	12	Not reported	Summary	No: only summary statistics reported
Nathan, 1998 202	V		Ø	V			Single	1	283	0	0	Prospective	Counts	No: only one patient group
Rosen, 1998 ²⁰¹		$\overline{\mathbf{Q}}$					Single	2	34	1	60	Prospective	Counts	Reported by authors
Scelsa, 1998 ²⁴³			Ø				Single	2	63	1	25	Prospective	Counts	Reported by authors
Seror, 1998 159			Ø				Single	1	85	1	80	Not reported	Counts	Reported by authors
Smith, 1998 ²⁴⁴			\square				Single	1	82	0	0	Prospective	Counts	Calculated by ECRI
Tan, 1998 ²⁰⁶			Ø		V		Single	1	64	1	56	Not reported	Summary	No: only summary statistics reported
Terzis, 1998 162			Ø				Single	1	72	1	43	Not reported	Counts	Reported by authors
Tetro, 1998 102	V	Ø	Ø			V	Single	1	64	1	50	Prospective	Counts	Reported by authors
Werner, 1998 ²⁰⁷	Ø		Ø	Ø			Multiple (>5)	1	727	0	0	Prospective	Counts	No: only one patient group
Wilson, 1998 ²⁴⁵			V	$\overline{\mathbf{A}}$			Single	1	23	1	14	Not reported	Summary	No: only summary statistics reported
Bak, 1997 ²⁴⁶				$\overline{\mathbf{A}}$	V		Single	1	20	0	0	Prospective	Counts	No: no control group
Brahme, 1997 199	V				\		Single	1	20	1	15	Prospective	Counts	Reported by authors

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?	
Bronson, 1997 163			Ø	Ø			Single	1	22	1	16	Prospective	Patient level	Calculated by ECRI	
Del Pino, 1997 104	Ø						Single	1	180	1	100	Prospective	Counts	Reported by authors	
Dellon, 1997 107	Ø	$\overline{\checkmark}$					Single	1	72	2	94	Not reported	Counts	No: inconsistent thresholds	
Franzblau, 1997 ²⁰⁸	Ø						Single	1	148	0	0	Prospective	Summary	No: only summary statistics reporte	
Guglielmo, 1997 ²⁴⁷			Ø	V			Single	1	198	1	69	Prospective	Summary	No: only summary statistics reported	
Gunnarsson, 1997 248			V				Single	1	100	0	0	Prospective	Counts	Reported by authors	
Horch, 1997 ²⁴⁹					V		Single	1	19	1	17	Not reported	Summary	No: only summary statistics reported	
Jeng, 1997 ²⁰⁹	Ø	V	$\overline{\square}$	V		V	Single	1	27	0	0	Prospective	Counts	Reported by authors	
Kaneko, 1997 ²⁵⁰			Ø	V			Single	1	15	3	66	Not reported	Summary	No: only summary statistics reported	
King, 1997 114		$\overline{\mathbf{A}}$					Single	1	29	1	100	Not reported	Summary	No: only summary statistics reported	
Pierre-Jerome, 1997 ²⁵¹			Ø		V		Single	1	27	1	28	Prospective	Summary	No: only summary statistics reported	
Radack, 1997 ²⁵²					V		Single	1	161	1	NR	Retrospective	Counts	Reported by authors	
Rosecrance, 1997				V		V	Single	1	28	1	25	Not reported	Summary	No: only summary statistics reported	
Simovic, 1997 182			Ø	V			Single	1	107	1	15	Retrospective	Counts	Reported by authors	
Werner, 1997 ²¹⁰			V	Ŋ			Single	2	108	0	0	Retrospective	Counts	No: incomplete reporting	
Andary, 1996 196			V	Ŋ			Single	1	81	1	17	Prospective	Counts	Reported by authors	
Atroshi, 1996 136			Ø				Single	1	36	2	60	Prospective	Counts	Reported by authors	
Bingham, 1996 ²¹¹	V		V	V			Single	1	1021	0	0	Prospective	Counts	No: only one patient group	
Checkosky, 1996 ²⁵⁴		7					Single	1	24	1	20	Not reported	Patient level	Reported by authors	
Cherniak, 1996 190		V	\square	V		V	Single	1	49	1	10	Not reported	Counts	Reported by authors	
Foresti, 1996 192			\square	$\overline{\mathbf{A}}$			Single	1	100	1	25	Prospective	Counts	Reported by authors	
Ghavanini, 1996 ²⁵⁵			\square	$\overline{\mathbf{A}}$			Single	1	50	1	50	Not reported	Summary	No: only summary statistics reported	
Kleindienst, 1996 ²⁵⁶					V		Single	1	55	1	18	Not reported	Counts	Reported by authors	
Murata, 1996 164	\square		\square	$\overline{\mathbf{A}}$			Single	1	27	1	19	Not reported	Counts	Calculated by ECRI	

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?	
Padua, 1996 165			$\overline{\checkmark}$	V			Single	1	43	1	36	Not reported	Counts	Reported by authors	
Pierre-Jerome, 1996 ²¹²	Ø		Ø		V		Single	1	24	1	19	Prospective	Summary	No: only summary statistics reported	
Britz, 1995 ²⁵⁷	V		V	V	V	Ø	Single	1	32	1	5	Prospective	Patient level	No: results not reported for controls	
De Smet, 1995 101	V						Single	2	50	2	55	Not reported	Counts	Reported by authors	
Gerr, 1995 118		$\overline{\checkmark}$					Single	2	60	1	59	Not reported	Counts	Reported by authors	
Glass, 1995 ²⁸	V		$\overline{\checkmark}$				Single	1	82	1	24	Not reported	Counts	Calculated by ECRI	
Golovchinsky, 1995			Ø	V		Ø	Single	1	571	0	0	Retrospective	Counts	Reported by authors	
Hamanaka, 1995 ²⁵⁹			V	V			Single	2	647	1	31	Retrospective	Counts	Calculated by ECRI	
Hansson, 1995 137			$\overline{\checkmark}$	$\overline{\mathbf{A}}$			Single	2	30	1	10	Not reported	Counts	Reported by authors	
Kothari, 1995 ²⁶⁰			V				Single	1	59	1	30	Not reported	Summary	No: only summary statistics reported	
Lang, 1995 109		$\overline{\mathbf{A}}$	$\overline{\checkmark}$			$\overline{\checkmark}$	Single	1	23	1	16	Prospective	Counts	Reported by authors	
Lesser, 1995 ²⁶¹			V	V		V	Single	1	45	1	20	Not reported	Counts	Reported by authors	
Nakamichi, 1995 ²⁶²					V		Single	1	15	1	15	Not reported	Patient level	Calculated by ECRI	
Seradge, 1995 ²⁶³						V	Single	1	72	1	21	Not reported	Summary	No: only summary statistics reported	
Seror, 1995 179			$\overline{\checkmark}$	$\overline{\mathbf{A}}$			Single	3	75	1	40	Not reported	Counts	Reported by authors	
Shafshak, 1995 ²⁶⁴			$\overline{\mathbf{A}}$			$\overline{\mathbf{A}}$	Single	2	36	2	36	Not reported	Counts	No: no diagnostic results reported	
Sheean, 1995 191			V	V			Single	1	49	1	NR	Not reported	Counts	Calculated by ECRI	
Tassler, 1995 115		$\overline{\mathbf{A}}$		$\overline{\mathbf{A}}$			Single	1	14	1	13	Retrospective	Counts	Reported by authors	
Valls-Sole, 1995 265			$\overline{\checkmark}$	$\overline{\mathbf{A}}$			Single	1	18	1	15	Prospective	Summary	No: only summary statistics reported	
Werner, 1995 ²¹³	V	V	V	V			Single	1	167	0	0	Not reported	Counts	Reported by authors	
Young, 1995 166	V	$\overline{\Delta}$	$\overline{\checkmark}$				Single	1	157	0	0	Prospective	Counts	No: only one patient group	
Clifford, 1994 ²⁶⁶			$\overline{\checkmark}$	$\overline{\mathbf{A}}$			Single	1	20	1	10	Not reported	Summary	No: only summary statistics reported	
Durkan, 1994 ²⁶⁷	V						Single	1	30	1	25	Not reported	Counts	Calculated by ECRI	
Franzblau, 1994 113	V	$\overline{\Delta}$	$\overline{\checkmark}$	$\overline{\square}$			Single	1	83	0	0	Prospective	Counts	Reported by authors	
Gerr, 1994 197	V	$\overline{\Delta}$					Single	2	NR	1	NR	Not reported	Counts	Reported by authors	

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?	
Kirschberg, 1994 ²¹⁴	Ø		$\overline{\mathbf{A}}$	V			Single	1	112	0	0	Retrospective	Counts	No: only one patient group	
Kuntzer, 1994 144			$\overline{\mathbf{A}}$	V		$\overline{\checkmark}$	Single	1	100	1	70	Prospective	Counts	Reported by authors	
Nathan, 1994 ²¹⁵	Ø		Ø	V			Multiple (<5)	2	417	0	0	Retrospective	Counts	No: no control subjects	
Nilsson, 1994 ²¹⁶			Ø				Single	3	175	0	0	Prospective	Counts	Reported by authors	
Para, 1994 103	Ø		Ø	V		V	Single	2	51	1	12	Not reported	Counts	Reported by authors	
Rossi, 1994 178			Ø	Ø			Single	1	62	1	27	Not reported	Counts	Reported by authors	
Werner, 1994 ²¹⁷	Ø	Ø	Ø	Ø			Single	1	130	0	0	Prospective	Counts	Calculated by ECRI	
Werner, 1994 111	Ø		Ø	V			Single	1	31	1	20	Not reported	Counts	Calculated by ECRI	
Eisen, 1993 193			Ø	Ø			Single	1	NR	1	NR	Not reported	Counts	Reported by authors	
Johnson, 1993 167	Ø		$\overline{\square}$				Single	1	184	0	0	Prospective	Summary	No: only summary statistics reported	
Nakamichi, 1993 ²⁶⁸					V		Single	1	128	0	0	Not reported	Counts	No: only one patient group	
Nathan, 1993 ²¹⁸	Ø		Ø	Ø			Single	2	1125	1	45	Prospective	Counts	Reported by authors	
Rodriquez, 1993 ²⁶⁹			Ø			Ø	Single	1	10	1	8	Prospective	Patient level	Calculated by ECRI	
Rosen, 1993 ²⁷⁰		V	Ø				Single	2	62	2	71	Not reported	Counts	Calculated by ECRI	
Rosén, 1993 138			Ø	Ø			Single	1	28	3	86	Not reported	Counts	Calculated by ECRI	
Uncini, 1993 160			$\overline{\square}$	$\overline{\mathbf{A}}$			Single	1	70	1	47	Not reported	Counts	Reported by authors	
Buchberger, 1992					V		Multiple (<5)	1	18	1	NR	Not reported	Counts	Reported by authors	
Grant, 1992 ²¹⁹		$\overline{\mathbf{A}}$	$\overline{\square}$				Single	1	22	1	47	Not reported	Counts	Calculated by ECRI	
Imaoka, 1992 ²⁷²			\square				Single	1	42	1	32	Not reported	Counts	Calculated by ECRI	
Kindstrand, 1992 ²⁷³						Ø	Single	1	94	1	127	Prospective	Patient level	Calculated by ECRI	
Preston, 1992 188			\square	V			Single	1	8	1	NR	Not reported	Counts	Calculated by ECRI	
Tchou, 1992 ²⁷⁴						Ø	Single	1	61	1	40	Not reported	Patient level	Reported by authors	
Buchberger, 1991					V		Single	1	25	1	14	Not reported	Summary	No: only summary statistics reported	

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?
Chang, 1991 145			\square				Single	1	43	1	40	Not reported	Counts	Calculated by ECRI
Durkan, 1991 155	V						Single	1	31	1	50	Not reported	Counts	Reported by authors
Jetzer, 1991 168	V	V				V	Single	3	323	1	284	Prospective	Counts	No: no control subjects
Katz, 1991 ²⁷⁶	V	V	V	V		Ø	Single	1	78	0	0	Not reported	Counts	Reported by authors
Lauritzen, 1991 185			V	V		Ø	Single	1	38	1	23	Not reported	Counts	Calculated by ECRI
Luchetti, 1991 169	Ø		Ø	Ø			Single	1	14	0	0	Retrospective	Patient level	No: only one patient group
Radwin, 1991 ¹¹⁶		$\overline{\mathbf{V}}$					Single	1	12	1	15	Not reported	Patient level	No: no diagnostic threshols used
Charles, 1990 170			Ø	\square			Single	1	158	2	90	Not reported	Counts	Reported by authors
De Krom, 1990 222	\square		V				Single	1	50	0	0	Prospective	Counts	Calculated by ECRI
Fitz, 1990 ²⁷⁷			\square				Single	1	36	1	44	Not reported	Counts	Calculated by ECRI
Gilliatt, 1990 278			V	V			Single	1	10	1	15	Not reported	Counts	Calculated by ECRI
MacDonell, 1990 90			Ø				Single	1	34	1	12	Not reported	Counts	Reported by authors
Merchut, 1990 279		V	V				Single	1	23	1	54	Not reported	Counts	Reported by authors
Palliyath, 1990 171			Ø				Single	1	10	1	11	Not reported	Summary	No: only summary statistics reported
Pease, 1990 177			V				Single	1	21	1	16	Not reported	Counts	Calculated by ECRI
Rojviroj, 1990 ²⁸⁰		V					Single	1	33	1	16	Prospective	Counts	Reported by authors
Tzeng, 1990 180			V				Single	1	84	1	50	Not reported	Counts	Calculated by ECRI
Uncini, 1990 135			V	V			Single	1	35	1	39	Not reported	Summary	No: only summary statistics reported
Winn, 1990 ²⁸¹		$\overline{\mathbf{A}}$					Single	2	61	0	0	Prospective	Summary	No: only summary statistics reported
Braun, 1989 ²⁸²	V	V					Single	1	40	0	0	Not reported	Counts	No: no diagnostic thresholds reported
Cioni, 1989 146			V	V			Single	1	307	1	54	Not reported	Counts	Reported by authors
Jackson, 1989 150			\square				Single	1	123	1	38	Not reported	Counts	Reported by authors
Meyers, 1989 ²⁸³						Ø	Single	1	14	1	19	Not reported	Counts	Calculated by ECRI
So, 1989 ¹⁷³			Ø			Ø	Single	1	22	2	35	Not reported	Counts	Reported by authors
Szabo, 1989 ²⁸⁴		$\overline{\Delta}$					Single	1	22	0	0	Not reported	Summary	No: only summary statistics reported
Uncini, 1989 161			$\overline{\square}$	$\overline{\mathbf{A}}$			Single	1	32	1	33	Not reported	Summary	No: only summary statistics reported
De Léan, 1988 ²⁸⁵			V				Single	1	150	0	0	Not reported	Counts	Calculated by ECRI

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?
Koris, 1988 198	$\overline{\Delta}$	$\overline{\mathbf{A}}$					Single	1	21	1	15	Prospective	Counts	Reported by authors
Molitor, 1988 110	Ø					V	Single	1	19	1	NR	Not reported	Counts	Calculated by ECRI
Mortier, 1988 ²⁸⁶							Single	1	116	1	102	Retrospective	Counts	Reported by authors
Pease, 1988 ²⁸⁷			Ø	$\overline{\mathbf{A}}$			Single	1	25	1	23	Not reported	Summary	No: only summary statistics reported
Carroll, 1987 ²⁸⁸			Ø	V			Single	1	101	1	50	Not reported	Counts	Reported by authors
Jessurun, 1987 ²⁸⁹					Ā		Multiple (<5)	1	24	1	10	Not reported	Summary	No: only summary statistics reported
Johnson, 1987 ²⁹⁰			Ø				Single	1	20	1	78	Not reported	Counts	Calculated by ECRI
Liang, 1987 ²⁹¹					V		Single	1	68	2	139	Not reported	Summary	No: only summary statistics reported
Macleod, 1987 ²⁹²	Ø		Ø				Single	1	111	1	125	Not reported	Summary	No: only summary statistics reported
Seror, 1987 156	V						Single	1	62	1	20	Not reported	Counts	Reported by authors
Borg, 1986 ²⁹³	V	V	V				Single	1	22	0	0	Not reported	Counts	Calculated by ECRI
Gellman, 1986 106	Ø		Ø				Single	1	NR	2	NR	Not reported	Counts	Reported by authors
Escobar, 1985 151			Ø				Single	1	23	1	55	Not reported	Counts	Calculated by ECRI
Kimura, 1985 ¹⁸⁹			Ø	V		V	Single	1	438	1	148	Not reported	Counts	Reported by authors
Mills, 1985 194			Ø	$\overline{\mathbf{A}}$			Single	1	47	2	49	Not reported	Counts	Calculated by ECRI
Borg, 1984 ²⁹⁴		V					Single	3	45	0	0	Prospective	Patient level	Calculated by ECRI
Pryse-Phillips, 1984	Ø						Single	1	212	4	184	Retrospective	Counts	Reported by authors
Satoh, 1984 ²⁹⁵	☑						Single	1	14	0	0	Retrospective	Patient level	No: only one patient group
Szabo, 1984 30	Ø	V					Single	1	20	0	0	Prospective	Counts	No: only one patient group
Goddard, 1983 ²⁹⁶			Ø				Single	1	24	1	49	Not reported	Counts	Calculated by ECRI
Kim, 1983 ¹⁹⁵			V	V			Single	1	39	1	33	Not reported	Counts	Reported by authors
Marin, 1983 139			Ø	V			Single	1	14	1	12	Not reported	Counts	Calculated by ECRI
Wongsam, 1983 172			Ø				Single	1	15	2	56	Not reported	Summary	No: only summary statistics reported
Johnson, 1981 ²⁹⁷			V	V			Single	1	18	1	37	Not reported	Summary	No: only summary statistics reported

Article	SGN	SEN	NCS	СМР	IMG	отн	Centers	CTS groups	CTS pts.	Neg. groups	Neg. subjects	Prospective or retrospective	Level of reporting	Could sensitivity & specificity be determined?	
Dekel, 1980 ²¹					V		Single	1	26	1	33	Prospective	Patient	No: could not extract 2 x 2 counts	
													level	from graph	
Messina, 1980 120			☑	\square			Single	1	40	1	40	Not reported	Counts	Reported by authors	
Gelmers, 1979 ²⁹	\square		\square	\square			Single	1	47	1	43	Not reported	Counts	Reported by authors	
Kimura, 1979 ¹⁴⁰			\square				Single	1	105	1	61	Not reported	Counts	Calculated by ECRI	
Schwartz, 1979 187							Single	1	20	1	10	Not reported	Counts	Calculated by ECRI	
Stewart, 1978 157	\square						Single	1	37	1	38	Not reported	Counts	Reported by authors	
Eisen, 1977 ²⁹⁸			Ø	V			Single	1	30	3	101	Not reported	Patient level	Calculated by ECRI	
Sedal, 1973 ²⁹⁹			\square				Single	1	214	1	34	Retrospective	Counts	Reported by authors	
Welch, 1973 ²²³		$\overline{\mathbf{Q}}$					Single	1	428	1	111	Not reported	Summary	No: only summary statistics reported	
Casey, 1972 300			Ø				Single	1	16	2	112	Not reported	Patient level	Calculated by ECRI	
Loong, 1972 141			V	Ŋ			Single	1	18	1	30	Not reported	Patient level	Calculated by ECRI	
Melvin, 1972 147			V				Single	1	17	1	24	Not reported	Counts	Calculated by ECRI	
Buchthal, 1971 301			Ø			$\overline{\Delta}$	Single	1	22	1	10	Not reported	Counts	Calculated by ECRI	
Loong, 1971 148			V	V			Single	1	15	1	30	Not reported	Patient level	Calculated by ECRI	
Plaja, 1971 142			Ø			V	Single	1	56	1	20	Retrospective	Counts	Reported by authors	

Table 45. Carpal Tunnel Syndrome-Patient Groups

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Finsen, 2001 224	CTS	Unspecified diagnosis	68	74	48	21	86				Yes
Mondelli, 2001 181	Normal	Healthy volunteers	19	NR	51.9	31	72				No
Mondelli, 2001 181	CTS	Unspecified diagnosis	20	80	52.8	35	75				No
Atroshi, 2000 225	CTS	Symptoms/ presented	262	57	52						No
Atroshi, 2000 225	Normal	Healthy volunteers	125	55	51						No
Bland, 2000 200	CTS	Complex objective standard	4690	65	57						No
Bland, 2000 200	CTS	Symptoms/ presented	8223	66	53	10	98				No
Bland, 2000 200	Normal	Other	3533	67	49						No
Cuturic, 2000 226	CTS	Unspecified diagnosis	19	0	43	29	62				No
Cuturic, 2000 226	Normal	Healthy volunteers	16	0	41	26	58				No
Kearns, 2000 ²⁰⁴	CTS	Workers at risk	45	4							Yes
Loscher, 2000 175	Normal	Healthy volunteers	87	NR	47	15	86				No
Loscher, 2000 175	CTS	Unspecified diagnosis		NR							No
Loscher, 2000 175	CTS	Other		NR							No
Montagna, 2000 ²²⁷	Cubital tunnel syndrome	Unspecified diagnosis	10	NR							No
Montagna, 2000 227	Normal	Healthy volunteers	15	NR							No
Montagna, 2000 ²²⁷	CTS	Unspecified diagnosis	30	NR							No
Nakamichi, 2000 ²²⁸	CTS	Simple nerve conduction	125	100	56	40	70				No
Nakamichi, 2000 ²²⁸	Normal	Healthy volunteers	200	NR	57	40	70				No
Raudino, 2000 ²²⁹	CTS	Complex objective standard	83	82	48.9	19	82	26.9	1	180	Yes
Resende, 2000 174	CTS	Unspecified diagnosis	20	NR							No
Resende, 2000 174	Normal	Healthy volunteers	20	NR		21	55				No
Resende, 2000 184	Normal	Healthy volunteers	20	100	36	20	54				No
Resende, 2000 184	CTS	Unspecified diagnosis	32	100	44	25	59				No
Sener, 2000 186	CTS	Symptoms/ presented	31	NR	46	26	70				Yes

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Sener, 2000 186	Normal	Healthy volunteers	21	NR	38	18	60				Yes
Seror, 2000 158	Normal	Healthy volunteers	20	75	43	20	67				No
Seror, 2000 158	CTS	Complex objective standard	20	75	47	32	76				No
Stalberg, 2000 ²³⁰	CTS	Symptoms/ presented	136	NR							No
Stalberg, 2000 ²³⁰	Normal	Healthy volunteers	32	NR		21	62				No
Weber, 2000 108	CTS	Symptoms/ presented	53	79	45						No
Weber, 2000 108	Normal	Healthy volunteers	26	85	37						No
Burke, 1999 ²³¹	CTS	Symptoms/ presented	186	NR							No
Atroshi, 1999 220	Normal	Other	2466	NR							No
Duncan, 1999 232	CTS	Complex objective standard	68	74	54						Yes
Duncan, 1999 232	CTS	Complex objective standard		NR							Yes
Duncan, 1999 232	Normal	Healthy volunteers	36	64	44						Yes
Kabiraj, 1999 ²³³	Normal	Healthy volunteers	38	50		20	79				No
Kabiraj, 1999 ²³³	CTS	Complex objective standard	31	68		28	85				No
Lee, 1999 ²³⁴	Normal	Healthy volunteers	28	54		22	47				No
Lee, 1999 ²³⁴	CTS	Unspecified diagnosis	50	74		32	81				No
Missere, 1999 ²⁰⁵	CTS	Workers at risk	45	0	37.7						No
Mongale, 1999 ²³⁵	Normal	Healthy volunteers	9	100	39	26	50				No
Mongale, 1999 ²³⁵	Normal	Healthy volunteers	7	0	39	27	58				No
Mongale, 1999 ²³⁵	CTS	Unspecified diagnosis	8	100	43	24	54				No
Murthy, 1999 143	CTS	Symptoms/ presented	84	NR							No
Murthy, 1999 143	Normal	Healthy volunteers	37	NR							No
Rudolfer, 1999 ²³⁶	CTS	Symptoms/ presented	937	NR							No
Sander, 1999 ²³⁷	Normal	Healthy volunteers	34	NR	41	26	71				No
Sander, 1999 ²³⁷	CTS	Complex objective standard	59	NR	49	29	73				No
Simovic, 1999 183	CTS	Other	12	NR							Yes
Simovic, 1999 183	Normal	Healthy volunteers	19	63	40	25	68				Yes
Simovic, 1999 183	CTS	Unspecified diagnosis	54	NR							Yes

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Szabo, 1999 152	Normal	Healthy volunteers	50	66		18	59				No
Szabo, 1999 152	CTS	Complex objective standard	50	76		20	73		2	240	No
Szabo, 1999 ¹⁵²	Unrelated disease	Other	50	80		28	72		0	180	No
Thonnard, 1999 117	CTS	Unspecified diagnosis	11	73	52						No
Thonnard, 1999 117	Normal	Healthy volunteers	11	73	53						No
Wang, 1999 ²³⁸	CTS	Complex objective standard	12	92	46	30	65				No
Wang, 1999 ²³⁸	Normal	Healthy volunteers	12	42	37	28	59				No
Aurora, 1998 ²³⁹	CTS	Symptoms/ presented	19	NR	52.8						No
Aurora, 1998 ²³⁹	Normal	Healthy volunteers	20	NR	32.9						No
Ferry, 1998 ²²¹	Normal	Other	648	56	46.9						No
Fertl, 1998 153	Normal	Healthy volunteers	20	60	42	25	77				No
Fertl, 1998 153	CTS	Symptoms/ presented	47	83	55.5	21	78				No
Gerr, 1998 31	Normal	Healthy volunteers	59	69	38.2						No
Gerr, 1998 31	CTS	Symptoms/ presented	60	72	46.6						No
Ghavanini, 1998 154	CTS	Complex objective standard	26	100	37	20	50	9	1	36	No
Ghavanini, 1998 154	CTS	Symptoms/ presented	74	81	40	20	50	15	1	60	No
Ghavanini, 1998 154	Normal	Healthy volunteers	58	76	36.7	20	50				No
Ghavanini, 1998 154	CTS	Complex objective standard	26	69	41	20	50	19.4	1	48	No
Ghavanini, 1998 154	CTS	Complex objective standard	22	73	42	30	50	19	4	60	No
Girlanda, 1998 149	CTS	Symptoms/ presented	41	93	39	24	65	48	1	180	Yes
Girlanda, 1998 149	Normal	Healthy volunteers	45	NR							Yes
Kabiraj, 1998 ²⁴⁰	CTS	Symptoms/ presented	72	NR							No
Kabiraj, 1998 ²⁴⁰	Normal	Healthy volunteers	65	45	39.8	20	75				No
Kleindienst, 1998 241	CTS	Complex objective standard		NR							No
Kleindienst, 1998 241	CTS	Other		NR							No
Kleindienst, 1998 241	CTS	Complex objective standard		NR							No
Kleindienst, 1998 241	CTS	Other		NR							No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Kleindienst, 1998 241	Normal	Healthy volunteers	18	83	51	43	59				No
Kleindienst, 1998 241	CTS	Complex objective standard		NR							No
Kleindienst, 1998 241	CTS	Unspecified diagnosis	77	82	54	22	79				No
Luchetti, 1998 ²⁴²	CTS	Unspecified diagnosis	39	79	31	26	45				No
Luchetti, 1998 ²⁴²	Normal	Healthy volunteers	12	83	27	24	36				No
Nathan, 1998 ²⁰²	CTS	Workers at risk	283	45	35.2						No
Rosen, 1998 ²⁰¹	Normal	Healthy volunteers	60	NR							No
Rosen, 1998 201	CTS	Workers at risk	20	5	46	26	65				No
Rosen, 1998 201	CTS	Unspecified diagnosis	14	100	53	33	78				No
Scelsa, 1998 ²⁴³	CTS	Other	21	48	46	10	69				No
Scelsa, 1998 ²⁴³	CTS	Unspecified diagnosis	42	76	50	25	85				No
Scelsa, 1998 ²⁴³	Normal	Healthy volunteers	25	44	42	23	63				No
Seror, 1998 159	CTS	Unspecified diagnosis	85	74	46	25	83				No
Seror, 1998 ¹⁵⁹	Normal	Healthy volunteers	80	64	42	22	68				No
Smith, 1998 ²⁴⁴	CTS	Symptoms/ presented	82	61	44	17	88	14	1	120	No
Tan, 1998 ²⁰⁶	CTS	Workers at risk	64	63		22	28				No
Tan, 1998 ²⁰⁶	Normal	Healthy volunteers	56	57		21	29				No
Terzis, 1998 162	CTS	Unspecified diagnosis	72	92	49.6						No
Terzis, 1998 162	Normal	Healthy volunteers	43	84	48.3						No
Tetro, 1998 102	Normal	Healthy volunteers	50	74	46.9	22	79				No
Tetro, 1998 102	CTS	Complex objective standard	64	64	49.3	21	83				No
Werner, 1998 ²⁰⁷	CTS	Workers at risk	727	54	42	25	69				Yes
Wilson, 1998 ²⁴⁵	Normal	Healthy volunteers	14	NR	52	33	76				No
Wilson, 1998 ²⁴⁵	CTS	Complex objective standard	23	NR	59	24	76				No
Bak, 1997 ²⁴⁶	CTS	Symptoms/ presented	20	55							Yes
Brahme, 1997 ¹⁹⁹	CTS	Unspecified diagnosis	20	90	37	21	61				No
Brahme, 1997 199	Normal	Healthy volunteers	15	47	35	22	60				No
Bronson, 1997 163	Normal	Other	16	56	29.5	21	44				Yes

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Bronson, 1997 163	CTS	Unspecified diagnosis	22	73	34.4	21	59				Yes
Del Pino, 1997 104	Normal	Healthy volunteers	100	78	49	37	67				No
Del Pino, 1997 104	CTS	Complex objective standard	180	81	50	16	84	37.9	1	216	No
Dellon, 1997 107	CTS	Unspecified diagnosis	72	NR							Yes
Dellon, 1997 ¹⁰⁷	Cubital tunnel syndrome	Unspecified diagnosis	42	NR							Yes
Dellon, 1997 107	Normal	Other	52	62							Yes
Franzblau, 1997 ²⁰⁸	CTS	Workers at risk	148	57	44.2						Yes
Guglielmo, 1997 ²⁴⁷	CTS	Symptoms/ presented	198	60	46	13	84				No
Guglielmo, 1997 ²⁴⁷	Normal	Healthy volunteers	69	57	40.3	20	86				No
Gunnarsson, 1997 ²⁴⁸	CTS	Symptoms/ presented	100	NR							No
Horch, 1997 ²⁴⁹	Normal	Healthy volunteers	17	71	43.4	24	58				No
Horch, 1997 ²⁴⁹	CTS	Simple nerve conduction	19	63	49.7	25	67				No
Jeng, 1997 ²⁰⁹	CTS	Workers at risk	27	52	40.2	23	57				No
Kaneko, 1997 ²⁵⁰	CTS	Unspecified diagnosis	15	87		40	54				Yes
Kaneko, 1997 ²⁵⁰	Normal	Healthy volunteers	46	22		25	45				Yes
Kaneko, 1997 ²⁵⁰	Cubital tunnel syndrome	Unspecified diagnosis	10	20		45	56				Yes
Kaneko, 1997 ²⁵⁰	Combined WRUEDs	Unspecified diagnosis	10	50		40	62				Yes
King, 1997 114	CTS	Unspecified diagnosis	29	62							No
King, 1997 114	Normal	Healthy volunteers	100	50							No
Pierre-Jerome, 1997 ²⁵¹	Normal	Healthy volunteers	28	100	45.1	26	67				No
Pierre-Jerome, 1997 ²⁵¹	CTS	Simple nerve conduction	27	100	51.9	16	78	36	12	72	No
Radack, 1997 252	CTS	Complex objective standard		NR							No
Radack, 1997 252	Normal	Unrelated disease		NR							No
Radack, 1997 252	CTS	Symptoms/ presented	161	53	37.4	13	86				No
Rosecrance, 1997 ²⁵³	CTS	Complex objective standard	20	70	41.5			a32			No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Rosecrance, 1997 ²⁵³	CTS	Complex objective standard	10	60	39.9			a14			No
Rosecrance, 1997 ²⁵³	Normal	Healthy volunteers	25	28	38.8						No
Rosecrance, 1997 ²⁵³	CTS	Complex objective standard	28	NR							No
Simovic, 1997 182	Normal	Healthy volunteers	15	NR		18	70				No
Simovic, 1997 182	CTS	Unspecified diagnosis	107	61	51	19	86				No
Werner, 1997 ²¹⁰	CTS	Workers at risk	59	64	40.1						No
Werner, 1997 ²¹⁰	Normal	Simple nerve conduction	49	67	41.7						No
Andary, 1996 196	Normal	Healthy volunteers	17	NR	36						No
Andary, 1996 196	CTS	Symptoms/ presented	81	NR	42						No
Atroshi, 1996 136	Normal	Healthy volunteers	30	57	36	25	62				Yes
Atroshi, 1996 136	CTS	Symptoms/ presented	36	69	52	20	87	a24	1	120	Yes
Atroshi, 1996 136	Normal	Healthy volunteers	30	70	40	19	65				Yes
Bingham, 1996 ²¹¹	CTS	Workers at risk	1021	29	30.1	17	60				No
Checkosky, 1996 ²⁵⁴	Normal	Healthy volunteers	10	70		25	44				No
Checkosky, 1996 ²⁵⁴	Normal	Healthy volunteers	20	75		25	67				No
Checkosky, 1996 ²⁵⁴	CTS	Symptoms/ presented	12	83		45	70				No
Checkosky, 1996 ²⁵⁴	CTS	Symptoms/ presented	24	79	46.7	27	70				No
Checkosky, 1996 ²⁵⁴	Normal	Healthy volunteers	10	80		46	67				No
Checkosky, 1996 ²⁵⁴	CTS	Symptoms/ presented	12	75		27	45				No
Cherniak, 1996 190	Normal	Healthy volunteers	10	70	37.1	26	52				No
Cherniak, 1996 190	CTS	Symptoms/ presented	49	33	43	19	71				No
Foresti, 1996 192	Normal	Healthy volunteers	25	28	42	18	69				Yes
Foresti, 1996 192	CTS	Symptoms/ presented	100	25	49	27	78				Yes
Ghavanini, 1996 ²⁵⁵	CTS	Unspecified diagnosis	50	82	38.6	27	59				Yes
Ghavanini, 1996 ²⁵⁵	Normal	Healthy volunteers	50	78	28.7	20	42				Yes
Kleindienst, 1996 ²⁵⁶	CTS	Other	55	82	54						No
Kleindienst, 1996 ²⁵⁶	Normal	Healthy volunteers	18	83	51						No
Murata, 1996 164	Normal	Healthy volunteers	19	100	24	19	31				Yes

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Murata, 1996 164	CTS	Workers at risk	27	100	25	19	37				Yes
Padua, 1996 165	Normal	Healthy volunteers	36	69	43.7	19	79				No
Padua, 1996 165	CTS	Symptoms/ presented	43	72	45.2	23	80	27	2	48	No
Pierre-Jerome, 1996 ²¹²	CTS	Workers at risk	24	100	44	26	59				Yes
Pierre-Jerome, 1996 ²¹²	Normal	Other	19	100	39.5	25	44				Yes
Britz, 1995 ²⁵⁷	CTS	Unspecified diagnosis	32	NR							No
Britz, 1995 ²⁵⁷	Normal	Healthy volunteers	0	NR							No
De Smet, 1995 101	CTS	Simple nerve conduction	10	70	42.8	22	53				No
De Smet, 1995 101	Normal	Healthy volunteers	46	100	51	34	76				No
De Smet, 1995 101	Normal	Other	9	100							No
De Smet, 1995 101	CTS	Symptoms/ presented	40	93	50.8	23	77				No
Gerr, 1995 ¹¹⁸	Symptomatic /normal NCS	Complex objective standard	30	60	43.9						No
Gerr, 1995 118	CTS	Complex objective standard	30	83	50.1						No
Gerr, 1995 118	Normal	Healthy volunteers	59	69	38.2						No
Glass, 1995 ²⁸	CTS	Symptoms/ presented	82	77		23	69				No
Glass, 1995 ²⁸	Normal	Contralateral arm	26	NR							No
Glass, 1995 28	Normal	Healthy volunteers	24	58		24	69				No
Golovchinsky, 1995 ²⁵⁸	Combined WRUEDs	Unspecified diagnosis	571	49	45.2	22	86				No
Hamanaka, 1995 ²⁵⁹	CTS	Unrelated disease	31	39	37.9	18	67				Yes
Hamanaka, 1995 ²⁵⁹	CTS	Unspecified diagnosis	647	61	53.9	21	87				Yes
Hansson, 1995 137	CTS	Symptoms/ presented	20	95	45	31	60	a 9	2	120	Yes
Hansson, 1995 137	Normal	Healthy volunteers	10	90	45	26	65	a9	2	120	Yes
Hansson, 1995 137	CTS	Complex objective standard	10	100	57	41	79	a 9	2	120	Yes
Kothari, 1995 ²⁶⁰	CTS	Symptoms/ presented	59	75	50	22	91				No
Kothari, 1995 ²⁶⁰	Normal	Healthy volunteers	30	70	36	21	70				No
Lang, 1995 109	CTS	Unspecified diagnosis	23	78	51.4			a36	12	420	No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Lang, 1995 109	Normal	Healthy volunteers	16	63	55						No
Lesser, 1995 ²⁶¹	Normal	Healthy volunteers	20	40	36	22	50				No
Lesser, 1995 ²⁶¹	CTS	Complex objective standard	45	73	52	27	79				No
Nakamichi, 1995 ²⁶²	CTS	Unspecified diagnosis	15	100	53.9	50	58				Yes
Nakamichi, 1995 ²⁶²	Normal	Healthy volunteers	15	100	54.4	50	58				Yes
Seradge, 1995 ²⁶³	CTS	Unspecified diagnosis	72	75	45.6	18	80				No
Seradge, 1995 ²⁶³	Normal	Unrelated disease	21	52		20	74				No
Seror, 1995 179	Normal	Healthy volunteers	40	70	53						No
Seror, 1995 179	CTS	Unspecified diagnosis	25	80	56						No
Seror, 1995 179	CTS	Unspecified diagnosis	25	84	52						No
Seror, 1995 179	CTS	Unspecified diagnosis	25	84	55						No
Shafshak, 1995 ²⁶⁴	CTS	Complex objective standard	25	52		22	40				Yes
Shafshak, 1995 ²⁶⁴	Other	Other	11	27		23	51				Yes
Shafshak, 1995 ²⁶⁴	Normal	Healthy volunteers	25	52	42	18	57				Yes
Shafshak, 1995 ²⁶⁴	CTS	Unspecified diagnosis	11	100		27	53				Yes
Sheean, 1995 191	CTS	Symptoms/ presented	49	71	56.2	29	84				No
Sheean, 1995 191	Normal	Healthy volunteers		NR		22	59				No
Tassler, 1995 115	Cubital tunnel syndrome	Unspecified diagnosis	13	NR							Yes
Tassler, 1995 115	CTS	Unspecified diagnosis	14	NR							Yes
Valls-Sole, 1995 265	CTS	Complex objective standard	18	100		34	53		6	144	No
Valls-Sole, 1995 265	Normal	Healthy volunteers	15	87		25	51				No
Werner, 1995 ²¹³	CTS	Workers at risk	167	NR							No
Young, 1995 166	CTS	Workers atrisk	157	82	39.9	20	64				No
Clifford, 1994 ²⁶⁶	CTS	Symptoms/ presented	20	100	43.1						No
Clifford, 1994 ²⁶⁶	Normal	Healthy volunteers	10	NR	26.7				_		No
Durkan, 1994 ²⁶⁷	CTS	Unspecified diagnosis	30	43	52	21	88				No
Durkan, 1994 ²⁶⁷	Normal	Healthy volunteers	25	NR						<u>-</u>	No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Franzblau, 1994 113	CTS	Workers at risk	83	53	33.8						No
Gerr, 1994 197	Normal	Healthy volunteers		NR	38						No
Gerr, 1994 197	CTS	Complex objective standard		NR	43						No
Gerr, 1994 197	CTS	Complex objective standard		NR	50						No
Kirschberg, 1994 ²¹⁴	CTS	Workers at risk	112	85	33.3						No
Kuntzer, 1994 144	Normal	Healthy volunteers	70	60	43	25	70				No
Kuntzer, 1994 144	CTS	Symptoms/ presented	100	80	51	26	85				No
Nathan, 1994 ²¹⁵	CTS	Workers at risk	316	47	40.4						No
Nathan, 1994 ²¹⁵	CTS	Workers at risk	101	26	38.6						No
Nilsson, 1994 ²¹⁶	CTS	Workers at risk	58	0	24.6						No
Nilsson, 1994 ²¹⁶	CTS	Workers at risk	61	0	37.4						No
Nilsson, 1994 ²¹⁶	CTS	Workers at risk	56	0	32.4						No
Para, 1994 103	CTS	Symptoms/ presented	24	71	51.6	26	62				No
Para, 1994 103	CTS	Symptoms/ presented	27	70	48.6	28	60				No
Para, 1994 103	Normal	Healthy volunteers	12	58	36.6	17	55				No
Rossi, 1994 178	CTS	Unspecified diagnosis	62	84	49.4	22	63				No
Rossi, 1994 178	Normal	Healthy volunteers	27	67	44.6	22	62				No
Werner, 1994 ²¹⁷	CTS	Workers at risk	130	56	34						No
Werner, 1994 111	CTS	Symptoms/ presented	31	NR							No
Werner, 1994 111	Normal	Healthy volunteers	20	NR							No
Eisen, 1993 193	CTS	Symptoms/ presented		NR							Yes
Eisen, 1993 193	Normal	Healthy volunteers		NR							Yes
Johnson, 1993 167	CTS	Workers at risk	184	NR							No
Nakamichi, 1993 ²⁶⁸	CTS	Unspecified diagnosis	128	74	54	33	86				No
Nathan, 1993 ²¹⁸	Normal	Healthy volunteers	45	47	19.8						No
Nathan, 1993 ²¹⁸	CTS	Workers at risk	388	63	39.4						No
Nathan, 1993 ²¹⁸	CTS	Workers at risk	737	28	42.4						No
Rodriquez, 1993 ²⁶⁹	Normal	Healthy volunteers	8	38	40.3	23	82				No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Rodriquez, 1993 ²⁶⁹	CTS	Unspecified diagnosis	10	80	43.8	22	83				No
Rosen, 1993 270	Normal	Healthy volunteers	21	48	33.6	20	50				No
Rosen, 1993 270	Normal	Healthy volunteers	50	0	41.5	27	63				No
Rosen, 1993 270	CTS	Symptoms/ presented	47	0	42.8	23	63				No
Rosen, 1993 ²⁷⁰	CTS	Symptoms/ presented	15	80	37.9	26	53				No
Rosén, 1993 138	Normal	Healthy volunteers	15	60	34	21	46				No
Rosén, 1993 138	Normal	Other	50	0	41.5	27	63				No
Rosén, 1993 138	CTS	Unspecified diagnosis	28	75	41	26	77				No
Rosén, 1993 138	Normal	Healthy volunteers	21	48	33.6	20	50				No
Uncini, 1993 160	Normal	Healthy volunteers	47	72	44.7	18	78				No
Uncini, 1993 160	CTS	Simple nerve conduction	70	86	49.3	26	78				No
Buchberger, 1992 271	Normal	Healthy volunteers		NR							No
Buchberger, 1992 271	CTS	Unspecified diagnosis	18	78	57	23	82				No
Grant, 1992 219	CTS	Complex objective standard	22	NR		22	71				Yes
Grant, 1992 219	Normal	Healthy volunteers	47	100		16	65				Yes
Grant, 1992 219	CTS	Workers at risk		NR							Yes
Grant, 1992 ²¹⁹	CTS	Symptoms/ presented		NR							Yes
Imaoka, 1992 ²⁷²	CTS	Unspecified diagnosis	42	79	50.3	20	76				Yes
Imaoka, 1992 ²⁷²	Normal	Healthy volunteers	32	59	49.2	24	76				Yes
Kindstrand, 1992 ²⁷³	Normal	Other	127	65	47.5	15	84				Yes
Kindstrand, 1992 273	CTS	Complex objective standard	94	73	50	19	95		1	121	Yes
Preston, 1992 188	Normal	Healthy volunteers		NR	31	18	50				Yes
Preston, 1992 188	CTS	Other	8	NR							Yes
Preston, 1992 188	CTS	Symptoms/ presented		NR	49	21	98				Yes
Tchou, 1992 ²⁷⁴	CTS	Unspecified diagnosis	61	NR							No
Tchou, 1992 ²⁷⁴	Normal	Healthy volunteers	40	50		22	45				No
Buchberger, 1991 ²⁷⁵	Normal	Healthy volunteers	14	64							No
Buchberger, 1991 ²⁷⁵	CTS	Symptoms/ presented	25	68	61	38	85				No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Chang, 1991 145	Normal	Healthy volunteers	40	NR	38.6	22	60				Yes
Chang, 1991 145	CTS	Symptoms/ presented	43	79	42.3	25	64				Yes
Durkan, 1991 155	CTS	Complex objective standard	31	74	45	22	79				No
Durkan, 1991 155	Normal	Healthy volunteers	50	NR							No
Jetzer, 1991 168	CTS	Workers at risk	100	NR							No
Jetzer, 1991 168	CTS	Workers at risk	284	NR							No
Jetzer, 1991 168	CTS	Workers at risk	39	NR							No
Jetzer, 1991 168	Normal	Healthy volunteers	284	NR							No
Katz, 1991 ²⁷⁶	CTS	Symptoms/ presented	78	63	43.4						Yes
Lauritzen, 1991 185	CTS	Unspecified diagnosis	38	68	53						Yes
Lauritzen, 1991 185	Normal	Healthy volunteers	23	NR							Yes
Luchetti, 1991 169	CTS	Unspecified diagnosis	14	93	41	21	64	31.3	2	120	Yes
Radwin, 1991 116	CTS	Unspecified diagnosis	12	58		29	60				No
Radwin, 1991 116	Normal	Healthy volunteers	15	NR	34.5	25	67				No
Charles, 1990 170	Other	Other	30	60	45.5	25	63				Yes
Charles, 1990 170	Normal	Healthy volunteers	60	80	45	23	76				Yes
Charles, 1990 170	CTS	Unspecified diagnosis	158	84	47.1	20	64				Yes
De Krom, 1990 222	Normal	Other	50	86							No
Fitz, 1990 ²⁷⁷	Normal	Healthy volunteers	44	NR	30	22	66				No
Fitz, 1990 ²⁷⁷	CTS	Complex objective standard	36	NR	52	25	88				No
Gilliatt, 1990 ²⁷⁸	CTS	Unspecified diagnosis	10	NR	44						No
Gilliatt, 1990 ²⁷⁸	Normal	Healthy volunteers	15	NR	42						No
MacDonell, 1990 90	CTS	Complex objective standard	34	NR	44	29	67				No
MacDonell, 1990 90	Normal	Healthy volunteers	12	NR	41	26	61				No
Merchut, 1990 ²⁷⁹	Normal	Healthy volunteers	54	NR	53						No
Merchut, 1990 ²⁷⁹	CTS	Symptoms/ presented	23	87	53	25	74				No
Palliyath, 1990 171	Normal	Healthy volunteers	11	NR	31						No
Palliyath, 1990 171	CTS	Unspecified diagnosis	10	NR	42	30	50				No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Pease, 1990 177	Normal	Healthy volunteers	16	NR		21	63				No
Pease, 1990 177	CTS	Unspecified diagnosis	21	NR							No
Rojviroj, 1990 ²⁸⁰	CTS	Complex objective standard	33	76	46.5	19	67	19	1	120	No
Rojviroj, 1990 ²⁸⁰	Normal	Healthy volunteers	16	25							No
Tzeng, 1990 180	CTS	Unspecified diagnosis	84	70	48	21	67				No
Tzeng, 1990 180	Normal	Healthy volunteers	50	56	46	20	65				No
Uncini, 1990 ¹³⁵	Normal	Healthy volunteers	39	NR	54	16	81				No
Uncini, 1990 ¹³⁵	CTS	Complex objective standard	35	80	49	28	68			8	No
Winn, 1990 ²⁸¹	CTS	Other	34	NR							No
Winn, 1990 ²⁸¹	CTS	Symptoms/ presented	27	NR							No
Braun, 1989 ²⁸²	CTS	Symptoms/ presented	40	80	38						Yes
Cioni, 1989 146	Normal	Healthy volunteers	54	65	38.3	18	68				No
Cioni, 1989 146	CTS	Symptoms/ presented	307	16	46.4	20	72				No
Jackson, 1989 ¹⁵⁰	CTS	Symptoms/ presented	123	82	52.6	21	85				Yes
Jackson, 1989 ¹⁵⁰	Normal	Healthy volunteers	38	76	42.2	21	66				Yes
Meyers, 1989 ²⁸³	Normal	Healthy volunteers	19	53	36	22	60				No
Meyers, 1989 ²⁸³	CTS	Unspecified diagnosis	14	64	51	36	68				No
So, 1989 ¹⁷³	Normal	Healthy volunteers	20	NR							No
So, 1989 ¹⁷³	Cubital tunnel syndrome	Unspecified diagnosis	15	NR							No
So, 1989 ¹⁷³	CTS	Unspecified diagnosis	22	NR							No
Szabo, 1989 ²⁸⁴	CTS	Unspecified diagnosis	22	73	51	24	79	29	7	120	Yes
Uncini, 1989 161	CTS	Symptoms/ presented	32	NR							No
Uncini, 1989 ¹⁶¹	Normal	Healthy volunteers	33	55		16	81				No
De Léan, 1988 ²⁸⁵	CTS	Simple signs/symptoms	150	73	47.6	18	84	31	1	144	Yes
Koris, 1988 198	CTS	Unspecified diagnosis	21	86	60	28	85		1	120	Yes
Koris, 1988 ¹⁹⁸	Normal	Healthy volunteers	15	NR		28	40				Yes
Molitor, 1988 110	CTS	Symptoms/ presented	19	NR							No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Molitor, 1988 110	Normal	Healthy volunteers		NR	49	23	79				No
Mortier, 1988 ²⁸⁶	CTS	Simple nerve conduction	116	67	49.2	20	82				No
Mortier, 1988 ²⁸⁶	Normal	Healthy volunteers	102	67	47.5	22	86				No
Pease, 1988 ²⁸⁷	Normal	Healthy volunteers	23	NR		21	62				No
Pease, 1988 ²⁸⁷	CTS	Unspecified diagnosis	25	NR							No
Carroll, 1987 ²⁸⁸	CTS	Symptoms/ presented	101	76	44.8	22	82				No
Carroll, 1987 ²⁸⁸	Normal	Healthy volunteers	50	48	46.7	16	82				No
Jessurun, 1987 ²⁸⁹	Normal	Healthy volunteers	10	50							No
Jessurun, 1987 ²⁸⁹	CTS	Unspecified diagnosis	24	88							No
Johnson, 1987 ²⁹⁰	Normal	Healthy volunteers	78	NR		20	79				Yes
Johnson, 1987 ²⁹⁰	CTS	Complex objective standard	20	NR							Yes
Liang, 1987 ²⁹¹	CTS	Other	10	100							No
Liang, 1987 ²⁹¹	CTS	Unspecified diagnosis	68	79	50	24	73				No
Liang, 1987 ²⁹¹	Normal	Contralateral arm	39	67							No
Liang, 1987 ²⁹¹	Normal	Healthy volunteers	100	50	45	20	69				No
Liang, 1987 ²⁹¹	CTS	Other	28	82							No
Liang, 1987 ²⁹¹	CTS	Other	20	90							No
Liang, 1987 ²⁹¹	CTS	Other	20	65							No
Liang, 1987 ²⁹¹	CTS	Other	58	76							No
Macleod, 1987 ²⁹²	CTS	Simple nerve conduction	111	NR							No
Macleod, 1987 ²⁹²	Normal	Healthy volunteers	26	58	39	17	63				No
Macleod, 1987 ²⁹²	Normal	Healthy volunteers	125	52	41	17	82				No
Seror, 1987 156	CTS	Symptoms/ presented	62	79	56.8	29	85				No
Seror, 1987 156	Normal	Healthy volunteers	20	75	55.7	34	79				No
Borg, 1986 ²⁹³	CTS	Symptoms/ presented	22	82	45.5			33			No
Gellman, 1986 106	CTS	Complex objective standard		NR							Yes
Gellman, 1986 106	Normal	Healthy volunteers		NR							Yes
Gellman, 1986 106	Other	Other		NR							Yes

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Escobar, 1985 151	CTS	Symptoms/ presented	23	70		22	55				Yes
Escobar, 1985 ¹⁵¹	Normal	Healthy volunteers	55	64		20	70				Yes
Kimura, 1985 189	Normal	Healthy volunteers	148	54	47.6	20	81				No
Kimura, 1985 189	CTS	Symptoms/ presented	438	65	51.4	18	85				No
Mills, 1985 194	CTS	Symptoms/ presented	47	77		29	74		0	60	No
Mills, 1985 194	Normal	Healthy volunteers	29	45		19	63				No
Mills, 1985 194	Normal	Other	20	50		19	75				No
Borg, 1984 ²⁹⁴	CTS	Unspecified diagnosis	21	NR							No
Borg, 1984 ²⁹⁴	CTS	Other	12	NR							No
Borg, 1984 ²⁹⁴	CTS	Unspecified diagnosis	12	NR							No
Pryse-Phillips, 1984 105	Other	Complex objective standard	44	NR							No
Pryse-Phillips, 1984 105	Cubital tunnel syndrome	Complex objective standard	67	NR							No
Pryse-Phillips, 1984 105	CTS	Complex objective standard	212	NR							No
Pryse-Phillips, 1984 105	Other	Complex objective standard	41	NR							No
Pryse-Phillips, 1984 105	Other	Complex objective standard	32	NR							No
Satoh, 1984 ²⁹⁵	CTS	Complex objective standard	14	100							No
Szabo, 1984 30	CTS	Unspecified diagnosis	20	50		32	81		2	180	No
Goddard, 1983 ²⁹⁶	CTS	Unspecified diagnosis	24	NR							No
Goddard, 1983 ²⁹⁶	Normal	Healthy volunteers	49	NR							No
Kim, 1983 ¹⁹⁵	Normal	Healthy volunteers	33	NR	41.3	20	68				No
Kim, 1983 ¹⁹⁵	CTS	Symptoms/ presented	39	NR							No
Marin, 1983 ¹³⁹	CTS	Unspecified diagnosis	14	86	49	23	79	13	1	24	No
Marin, 1983 ¹³⁹	Normal	Healthy volunteers	12	42	30	22	48				No
Wongsam, 1983 172	DM with peripheral neuropathy	Unrelated disease	6	NR							No
Wongsam, 1983 172	CTS	Symptoms/ presented	15	NR							No

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Wongsam, 1983 172	Normal	Healthy volunteers	50	56		20	68				No
Johnson, 1981 ²⁹⁷	CTS	Unspecified diagnosis	18	NR							No
Johnson, 1981 ²⁹⁷	Normal	Healthy volunteers	37	49							No
Dekel, 1980 ²¹	Normal	Healthy volunteers	33	58	40.3						No
Dekel, 1980 ²¹	CTS	Unspecified diagnosis	26	100							No
Messina, 1980 120	CTS	Symptoms/ presented	40	NR	45.1	19	67				No
Messina, 1980 120	Normal	Healthy volunteers	40	NR	47.5						No
Gelmers, 1979 ²⁹	Normal	Healthy volunteers	43	79	54	26	74				No
Gelmers, 1979 ²⁹	CTS	Complex objective standard	47	81	57	29	78				No
Kimura, 1979 ¹⁴⁰	CTS	Unspecified diagnosis	105	70	48	20	78				No
Kimura, 1979 ¹⁴⁰	Normal	Unrelated disease	61	57	43	15	50				No
Schwartz, 1979 187	CTS	Symptoms/ presented	20	85	52	27	77				No
Schwartz, 1979 187	Normal	Healthy volunteers	10	100		20	28				No
Stewart, 1978 157	CTS	Complex objective standard	37	81	55	36	84				Yes
Stewart, 1978 157	Normal	Healthy volunteers	38	79	53	30	84				Yes
Eisen, 1977 ²⁹⁸	Cubital tunnel syndrome	Complex objective standard	18	NR	51.7	26	65				No
Eisen, 1977 ²⁹⁸	Normal	Healthy volunteers	60	NR	41.5	11	74				No
Eisen, 1977 ²⁹⁸	Combined WRUEDs	Other	23	NR	50	7	68				No
Eisen, 1977 ²⁹⁸	CTS	Complex objective standard	30	NR	56.1	21	76				No
Sedal, 1973 ²⁹⁹	Normal	Healthy volunteers	34	NR	47	18	77				Yes
Sedal, 1973 ²⁹⁹	CTS	Complex objective standard	214	56	54	19	87				Yes
Welch, 1973 ²²³	Other	Other	111	NR							No
Welch, 1973 ²²³	Combined WRUEDs	Workers at risk	428	81							No
Casey, 1972 300	CTS	Unspecified diagnosis	16	94	55.9	35	70				Yes
Casey, 1972 300	Other	Other	18	33	53.5	30	77	178	72	444	Yes

Article	Disorder type	Patient selection	N patients	% female	Mean age	Age of youngest	Age of oldest	Duration of condition before treatment (months)	Shortest duration (months)	Longest duration (months)	Are patient comorbidities reported?
Casey, 1972 300	Normal	Healthy volunteers	94	NR	51	20	80				Yes
Loong, 1972 141	Normal	Healthy volunteers	30	100		30	60				No
Loong, 1972 141	CTS	Unspecified diagnosis	18	100	43.7	31	60	12.7	1	48	No
Melvin, 1972 147	CTS	Symptoms/ presented	17	NR							No
Melvin, 1972 147	Normal	Healthy volunteers	24	NR							No
Buchthal, 1971 301	Normal	Healthy volunteers	10	50		32	57				No
Buchthal, 1971 301	CTS	Other	22	73		29	67			360	No
Loong, 1971 148	Normal	Healthy volunteers	30	100		30	60				Yes
Loong, 1971 148	CTS	Symptoms/ presented	15	100		31	60	7.6	1	24	Yes
Plaja, 1971 142	Normal	Healthy volunteers	20	NR							No
Plaja, 1971 142	CTS	Unspecified diagnosis	56	NR							No

^aReported median age instead of mean age CTS—Carpal tunnel syndrome DM—Diabetes mellitus

Table 46. Carpal Tunnel Syndrome—Reported Inclusion and Exclusion Criteria

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Finsen, 2001 ²²⁴	Positive clinical diagnosis of carpal tunnel syndrome	Patients for whom the clinical diagnosis was considered equivocal. If more than one hand was treated, only the first was included.
Mondelli, 2001 181	Idiopathic CTS with reduction of distal conduction velocity of the median nerve. Unilateral CTS.	None reported
Atroshi, 2000 ²²⁵	Respondents to a random survey who reported numbness and/or tingling in at least two radial fingers at least twice a week for previous four weeks	Previous CTS surgery, resolution of symptoms, symptoms not consistent with CTS, unwilling to take test
Bland, 2000 ²⁰⁰	All patients in county referred for NCS with suspected CTS, also patients with other referrals who then had a positive NCS	None (authors report 100% inclusion)
Cuturic, 2000 ²²⁶	Sensory symptoms and abnormal NCS, limited to mild or moderate disease	Certain EMG abnormalities (authors do not specify that these were in fact exclusion criteriajust that no patients had them)
Kearns, 2000 ²⁰⁴	Pork processing employees who had worked for at least 2 months.	Pre-existing CTS or diabetes.
Loscher, 2000 175	Referred to the laboratory for neurophysiological assessment of median nerve	Traumatic nerve lesions
Montagna, 2000 ²²⁷	Diagnosed with carpal tunnel syndrome or cubital tunnel syndrome.	None reported
Nakamichi, 2000 ²²⁸	DML >4.2 ms and SCV >45 m/s	None reported
Raudino, 2000 ²²⁹	Referred to lab. All were complaining of discomfort, paresthesias, or weakness in the territory of the median nerve occurring especially at night or after repetitive actions and relieved by changes in posture or shaking hands. Abnormal nerve conduction test as defined by one of the following three abnormalities: 1) DML >4 ms; 2) antidromic DSL to index finger >3 ms; wrist to-palm sensory latency >1.8 ms for patients <45 years old or >2 ms for patients older than 45.	Metabolic diseases, radiculopathies, polyneuropathies, concomitant pathologies.
Resende, 2000 ¹⁸⁴	Clinical diagnosis of carpal tunnel syndrome and abnormal conventional motor and sensory conduction studies	None reported
Resende, 2000 ¹⁷⁴	Diagnosed with carpal tunnel syndrome by clinical and electrophysiological methods with conventional techniques. Normal bilateral sensory conduction studies of the ulnar nerve.	None reported
Sener, 2000 ¹⁸⁶	Symptoms and clinical signs suggesting carpal tunnel syndrome.	Peripheral nerve dysfunction or peripheral neuropathy other than CTS
Seror, 2000 158	Diagnosis of mild CTS	None reported
Stalberg, 2000 ²³⁰	Patients referred to the lab with the presumptive diagnosis of carpal tunnel syndrome.	None reported
Weber, 2000 108	Suspected of having carpal tunnel syndrome.	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Atroshi, 1999 220	Randomly selected from the population of Sweden.	Did not respond to mailed questionnaire, did not attend clinical exam, previous carpal tunnel surgery, declined nerve conduction testing, neurologic disease
Burke, 1999 231	Referred for splinting	None reported
Duncan, 1999 ²³²	Positive NCS (decreased median SCV or prolonged DML) or two physicians agreeing that the symptoms and history are consistent with CTS. Did not give specific criteria for either.	Previous surgery or anatomic variation in the median nerve
Kabiraj, 1999 ²³³	DML >4.02 m/sec [sic] (mean + 2 SD), MCV <47.57 m/s (mean – 2 SD), CMAP decreased by 1 SD, prolonged or absent median sensory action potential. Painful paresthesia with night worsening, appropriate distribution, thenar weakness, positive Tinel, positive Phalen.	None reported
Lee, 1999 ²³⁴	Clinical diagnosis of CTS.	None reported
Missere, 1999 ²⁰⁵	Male workers in a meat processing plant	None reported
Mongale, 1999 235	Diagnosed with carpal tunnel syndrome via NCS.	None reported
Murthy, 1999 143	Referred for electrodiagnostic evaluation for paresthesia	None reported
Rudolfer, 1999 236	Patients in database referred to electromyographer.	Non-CTS abnormality.
Sander, 1999 ²³⁷	Both clinical and electrophysiological diagnosis of carpal tunnel. 1) Clinical: Two or more of the following primary symptoms in a median nerve distribution: numbness, tingling, clumsiness, or nocturnal symptom exacerbation. If only one of these symptoms was present, two of the following secondary symptoms were required: burning/cold, tightness, sore/ache/discomfort, or puffiness. 2) Electrodiagnostic confirmation: one of the following three abnormalities: A) an absent median palm-wrist mixed nerve action potential latency. B) a median palm-wrist mixed nerve action potential latency >1.7ms, C) if this same latency exceed the ipsilateral ulnar palm-wrist latency by more than 0.3ms.	Carpal tunnel patients: excluded if a history or physical exam suggestive of a neuromuscular disorder other than carpal tunnel syndrome.
Simovic, 1999 ¹⁸³	Referred to laboratory with hand or arm complaints including but not limited to numbness, tingling, or pain	Diabetes or the clinical or electrophysiological suggestion of a concomitant peripheral nerve disorder
Szabo, 1999 152	Diagnosed CTS	None reported
Thonnard, 1999 ¹¹⁷	Severe CTS: small or absent sensory amplitude, DSL and DML >5 ms, and evidence of denervation in APB	Other (non-CTS) electrodiagnostic abnormalities

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Wang, 1999 ²³⁸	Symptoms and at least 2 of the following 5 NCS criteria: 1) DML >4.2 ms 2) DSL to index >3.5 ms 3) Difference between median and ulnar mixed nerve latencies = 0.4 ms 4) Difference between median and ulnar sensory latency to ring finger = 0.5 ms 5) Difference between median motor latency to 2nd lumbrical and ulnar motor latency to first palmar interosseous = 0.5 ms	Additional neuromuscular disease, polyneuropathy, cervical radiculopathy, severe CTS, atypical histories.
Aurora, 1998 ²³⁹	Referred to lab with clinically definite carpal tunnel syndrome.	None reported
Ferry, 1998 ²²¹	All participants were registered to receive primary care at a local general practice.	None reported
Fertl, 1998 ¹⁵³	Referred with pain	Polyneuropathy, ulnar nerve lesion, radiculopathy, arthropathy
Gerr, 1998 ³¹	Any patient 18-70 years old with symptoms of pain, weakness, numbness, or tingling in the cutaneous distribution of the median nerve	Electrophysiological tests positive for a disorder other than CTS.
Ghavanini, 1998 154	Symptoms of CTS	Conditions other than CTS
Girlanda, 1998 ¹⁴⁹	Symptomatic hands with clinical evidence of idiopathic CTS. Examples of symptoms: nocturnal or activity-related pain and paresthesia in the hand, Phalen's, hypaesthesia limited to the distribution of the median nerve. Mild CTS required: No weakness or muscle atrophy present, DML in all patients was never slower than 4.0 ms which represented 2.5 SD below mean of controls in this laboratory.	Known causes of entrapment neuropathies or systemic diseases. Cervical radiculopathy, brachial plexopathy, thoracic outlet syndrome, multi-polyneuropathies.
Kabiraj, 1998 ²⁴⁰	Patients had the following symptoms and signs: history of pain, numbness, paresthesia, nocturnal awakening due to pain and weakness with or without atrophy, decreased sensations, Tinel's signs and wrist flexion Phalen's signs	Evidence of peripheral neuropathy other than median nerve dysfunction
Kleindienst, 1998 241	Clinical diagnosis of CTS	None reported
Luchetti, 1998 ²⁴²	Idiopathic CTS, defined as night pain and/or paresthesia, and median nerve sensory deficits. Motor deficits not required.	Diabetes, uremia, polyneuropathy, history of wrist trauma
Nathan, 1998 ²⁰²	Industrial workers in four industries: steel mill workers, food processors, electronics workers, and plastics workers.	Previous carpal tunnel release surgery.
Rosen, 1998 ²⁰¹	Carpal tunnel patients: Clinically diagnosed. Vibration-exposure patients Symptomatic, with exposure to hand-held vibrating tools.	None reported
Scelsa, 1998 ²⁴³	Clinically definite CTS as defined by: symptoms of numbness, paresthesia or pain in median nerve distribution and at least one of the following: hand clumsiness, nocturnal hand symptoms, sensory loss, weakness on exam in an appropriate median nerve distribution. Normal ulnar sensory and motor conduction studies	Cervical radicular pain or objective signs of cervical radiculopathy, or clinical evidence of polyneuropathy, or electrophysiological evidence of ulnar neuropathy

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Seror, 1998 159	Intermittent symptoms of burning, tingling, and paresthesia in the radial digits especially at night or upon awakening. Also patients had normal classical electrodiagnostic tests, i.e., DML to APB <4ms and palm-to-wrist orthodromic sensory conduction velocity >45m/s	None reported
Smith, 1998 ²⁴⁴	Referred with suspected CTS	None reported
Tan, 1998 ²⁰⁶	Working as carpet weaver	None reported
Terzis, 1998 ¹⁶²	CTS patients: Median distal motor latency required to be less than 4.2 ms. 18 months after the study, confirmation of CTS by sensory nerve latency on either digit 2 or digit 3 of >3ms.	Any history of peripheral nerve problems. Any other pathology, screened out by ulnar nerve and palmar stimulation studies
Tetro, 1998 ¹⁰²	CTS symptoms including median distribution of pain and paresthesia. Positive NCS including abnormal DML or DSL or DML 1.0 ms more than contralateral or DSL 0.5 ms more than contralateral	Proximal entrapment symptoms, thoracic outlet syndrome, acute CTS, paralysis, negative NCS (n = 7)
Werner, 1998 ²⁰⁷	Workers were selected to be representative of a range of jobs typically found in contemporary manufacturing and clerical sites.	None reported
Wilson, 1998 ²⁴⁵	Presence of carpal tunnel syndrome	History of significant hand trauma, or peripheral neuropathy, or radiculopathy, or Martin-Gruber anastomosis
Bak, 1997 ²⁴⁶	Suspected CTS	Diabetes, severe renal disease, pregnancy within the last year, previously treated CTS, contraindications to MRI, polyneuropathy.
Brahme, 1997 ¹⁹⁹	Diagnosed by hand surgeon with work-related dynamic carpal tunnel syndrome (indicating that symptoms only occurred during repetitive motion).	None reported
Bronson, 1997 ¹⁶³	Patients: Pre-surgery, DML <4 ms, normal needle EMG of APB. Included in this group based on traditional clinical indications, as judged by physicians. Controls: positive Tinel's sign, but no symptoms. Negative on standard sensory and motor nerve conduction tests.	Diabetes, rheumatoid arthritis, hypothyroidism, cervical spine disease, pregnancy, cervical radiculopathy.
Del Pino, 1997 ¹⁰⁴	All of the following three criteria for diagnosis of CTS: 1) Symptoms of CTS, consisting of pain predominantly at night, paresthesias and dysaesthesias, numbness, sensory deficit in the territory of the median nerve, and weakness of the APB; 2) Abnormal sensitivity in the median nerve distribution compared to the ulnar territory of the same hand and/or cutaneous territory of the contralateral median nerve in cases of unilateral involvement; 3) Complete relief of pain and paresthesias within 15 days of open surgical release of the carpal tunnel.	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Dellon, 1997 ¹⁰⁷	Already diagnosed with either carpal tunnel syndrome or cubital tunnel syndrome. Diagnosis was based on the clinical history and physical examination, which included positive provocative testing, positive Tinel's sign at the wrist or elbow, abnormal tuning fork perception.	Cervical radiculopathy, diabetes, thoracic outlet syndrome, thyroid disease, collagen vascular disease, using narcotics or antidepressants.
Franzblau, 1997 ²⁰⁸	At least 6 months' tenure in jobs at a spark plug manufacturing plant	None reported
Guglielmo, 1997 ²⁴⁷	Typical signs and symptoms of carpal tunnel syndrome (based on American Academy of Neurology Quality Standards Subcommittee)	None reported
Gunnarsson, 1997	Referred to lab with suspected CTS	Neuropathies
Horch, 1997 ²⁴⁹	Surgical candidates with symptoms of CTS and median motor latency >4 ms	None reported
Jeng, 1997 ²⁰⁹	Volunteers from food processing plant.	History of peripheral neuropathy, fractures, severe burns, arthritis, diabetes, carpal tunnel surgery
Kaneko, 1997 ²⁵⁰	Group 01: Coexisting entrapment neuropathy and cervical cord compression demonstrated by MRI. Group 02: Diagnosed with carpal tunnel syndrome. Group 03: Diagnosed with cubital tunnel syndrome. Group 04: Control group, no subjective symptoms or neurologic findings associated with peripheral or central lesions.	None reported
King, 1997 114	CTS as confirmed by EMG or NCS. New referrals.	None reported
Pierre-Jerome, 1997 ²⁵¹	Typical signs and symptoms, DML >4.5 ms or sensory velocity <45 m/s	Previous surgery, comorbidity with "somatic connective tissue diseases" (radiculopathy?), alcoholism
Radack, 1997 252	All wrist MRI examinations, regardless of indication	None
Rosecrance, 1997 253	Recent (within two weeks) numbness and tingling, or one of those plus any two of: burning/cold, tightness, pain, symptoms worsening at night. Must have involved median nerve distribution (thumb to medial aspect of ring finger).	Disorders with similar presentation to CTS.
Simovic, 1997 ¹⁸²	1) Referral to laboratory for possible carpal tunnel syndrome; and 2) Completion of a median motor study including distal and proximal stimulation, sensory antidromic median conduction to the index finger, and mixed nerve median and ulnar conduction studies with palmar stimulation	Clinical symptoms or signs of other peripheral nerve disorders of the same limb. 2) Diabetes mellitus Insufficient chart data
Werner, 1997 ²¹⁰	DSL prolonged by 0.5 ms or more, but asymptomatic	None reported
Andary, 1996 ¹⁹⁶	Referred to lab because of pain or numbness in the hand and wrist with histories and physical exam consistent with the possible diagnosis of CTS. Median antidromic sensory latency to index finger was required to be <4.0 ms to rule out "clear cut" CTS. Other nerve conduction tests (unspecified), however, were required to be positive.	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Atroshi, 1996 136	Symptoms and signs consistent with carpal tunnel syndrome. Unsuccessful prior nonoperative treatment.	None reported
Bingham, 1996 ²¹¹	All new applicants who had been offered jobs at meat packing, plastics assembly, food processing, furniture manufacturing, or grocery warehousing in a 17 county area in the southeastern US over an 18 month period. Applicants had worked for an average of 4.4 years in various settings.	None reported
Checkosky, 1996 ²⁵⁴	Physician-diagnosed CTS	None reported
Cherniak, 1996 190	Referred to lab.	None reported
Foresti, 1996 192	Patients with suspected carpal tunnel referred to the laboratory	Other pathologies potentially causing polyneuropathy such as diabetes, iperuremia, acromegaly, etc.
Ghavanini, 1996 ²⁵⁵	Paresthesia or numbness in fingers, and nocturnal hand pain or paresthesia, and excessive hand sweating or coldness, and positive Tinel sign or Phalen sign.	Diabetes, rheumatoid arthritis, thyroid dysfunction, history of trauma to neck or hands, cervical spondylosis, pregnancy, hand edema, obesity
Kleindienst, 1996 ²⁵⁶	Pre-operative	None reported
Murata, 1996 164	Data entry operators.	None of the patients complained of nocturnal awakening with paresthesia or pain in hands, none had positive Tinel's sign or positive Phalen's sign. Also excluded prior pregnancy, occupational exposure to neurotoxic substances, endocrine disorders, neurological disorders, diabetes, acromegaly, myxedema, lupus, amyloidosis, rheumatoid arthritis, alcoholic dependency, hand injury, forearm injury.
Padua, 1996 ¹⁶⁵	Paresthesia, pain, hypotrophy of thenar eminence	Other neuropathies or signs of severe CTS (i.e., absence of SNAP at wrist).
Pierre-Jerome,	Cleaners: Worked for at least three consecutive	Systemic diseases and psychiatric
1996 ²¹² Britz, 1995 ²⁵⁷	years and at least 19 hours a week. select group of patients who had been clinically diagnosed as having CTS	disorders including alcoholism. None reported
De Smet, 1995 101	Presented as surgical candidate	None reported
Gerr, 1995 ¹¹⁸	Age 18-70 with any hand symptoms	None reported
Glass, 1995 ²⁸	CTS symptoms	None reported
Golovchinsky, 1995	Referred to lab with complaints of neck pain and/or pain, numbness, or weakness in upper extremities.	Obvious injuries of the wrist, diabetes, hypothyroidism, renal failure.
Hamanaka, 1995 ²⁵⁹	Clinical diagnosis of CTS based on symptoms, sensory disturbance of the median nerve distribution area, Tinel's sign, Phalen's sign, manual muscle testing, and APB atrophy. Carpal canal pressure in resting position >15 mm Hg or carpal canal pressure in power active flex >135 mmHg.	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Hansson, 1995 137	Typical history (defined by sensory or motor symptoms like intermittent paresthesias, numbness, pain and weakness in the domain of the median nerve)	Diabetes, polyneuropathy, or rheumatic disease
Kothari, 1995 ²⁶⁰	Clinical diagnosis of CTS, including arm or wrist pain, paresthesia or other median distribution symptoms, weakness, Tinel's, or Phalen's and positive NCS	Signs or symptoms of neuropathy
Lang, 1995 ¹⁰⁹	CTS-typical signs and symptoms; 2) DML >4.5 ms or orthodromic SCV palm-to-wrist <45 m/s 3) planned surgical treatment	Previous surgery on the same hand
Lesser, 1995 ²⁶¹	Typical signs and symptoms of carpal tunnel syndrome, AND one or more of the following: 1) median distal motor latency >4.4ms, 2)median sensory antidromic latency to peak >3.5ms, 3) median sensory palm to wrist latency at least 0.4ms longer than that latency for the analogous segment of the ulnar nerve.	Peripheral neuropathy or multiple mononeuropathy
Nakamichi, 1995 ²⁶²	Clinical and electrophysiological diagnosis of bilateral CTS. Clinical evaluation included the presence of typical sensory symptoms, Phalen's test, two-point discrimination, muscle testing, and thenar atrophy. Electrophysiological criteria were either DML >4.2 ms or SCV <45 m/s.	Rheumatoid arthritis, chronic renal failure under hemodialysis, endocrine or metabolic disorders including diabetes, gout, amyloidosis, or hypothyroidism, Colles fracture, ganglion, calcium deposition, and osteoarthritis.
Seradge, 1995 ²⁶³	None reported	None reported
Seror, 1995 ¹⁷⁹	Referred to lab based on a clinical diagnosis of carpal tunnel syndrome: Intermitted paresthesia, numbness, tingling, or hypoesthesia in the median nerve distribution, with nocturnal aggravation, with or without pain in the hand, wrist, and forearm, and rarely for thenar muscle atrophy.	None reported
Shafshak, 1995 ²⁶⁴	Group 001: Positive Phalen's, positive Tinel's, DSL >4 ms, DML >4.7 ms, but normal ulnar nerve conduction studies Group 002: Definite polyneuropathy, DML >4.7 ms, slowed MCV at the forearm. Group 003: Severe unilateral CTS based on clinical findings, and unobtainable DML and DSL, but normal ulnar nerve conduction.	None reported
Sheean, 1995 191	Referred to lab based on suspected CTS.	None reported
Tassler, 1995 115	Symptomatic patients who had been diagnosed, had not been cured by nonoperative methods, and later received surgery for the condition.	Diabetes, alcoholism, other toxicity.
Valls-Sole, 1995 ²⁶⁵	Referred to lab, and all of the following:1) Slowing of MCV in wrist to palm and normal DML to thenar and normal CV elbow to wrist2) Normal CMAP amplitude from wrist or elbow stimulation3) Slow median SCV from palm to wrist, but no reduced SNAP amplitude4) Normal ulnar SCV5) No significant limitation of joint movement because of pain, skin or joint diseases or fat.	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Werner, 1995 ²¹³	Employees at an automobile parts manufacturing plant and a furniture assembly plant in southern Michigan.	None reported
Young, 1995 166	Workers at a poultry processing plant.	None reported
Clifford, 1994 ²⁶⁶	Referred to lab from family physicians, rheumatologists, and neurologists. Sy mptoms of CTS (e.g. pain, numbness, tingling). Screening history and physical exam to ensure the referring diagnosis of CTS was warranted.	Peripheral neuropathy, or obvious entrapment other than the median nerve.
Durkan, 1994 ²⁶⁷	Symptoms of CTS, particularly in median nerve distribution	None reported
Franzblau, 1994 ¹¹³	Full-time employees of an automobile parts manufacturing plant which had reported problems with upper extremity cumulative trauma disorders.	None reported
Gerr, 1994 ¹⁹⁷	Referred to lab, age 18-70 with symptoms of pain, weakness, numbness, or tingling that involved either hand.	None reported
Kirschberg, 1994 ²¹⁴	Employees in repetitive jobs in the poultry industry who were referred to a neurologist with pain, numbness, or tingling.	None reported
Kuntzer, 1994 ¹⁴⁴	If patient reported a combination of hand and arm symptoms suggestive of CTS, with numbness, tingling, pins and needles, "sleeping" of the hands and fingers, nocturnal symptoms or clumsiness, weakness, puffiness, swelling, tightness, joint pain or aching of the hand or fingers.	Patients: Two were excluded due to absent distal reflexes in the lower extremities. Controls: Two were excluded due to presence of symptoms of CTS, or pregnancy.
Nathan, 1994 ²¹⁵	Japanese furniture factory workers. American workers from four industries.	None reported
Nilsson, 1994 ²¹⁶	Currently working as a platers, truck assembler, or office worker. Male, age <54, randomly selected from larger groups for participation in the study. Platers were required to be currently exposed to vibration, and were selected for nerve conduction based on consecutive cases.	None reported
Para, 1994 ¹⁰³	Paresthetic CTS: Has CTS, has normal distal motor latency. Slight CTS: Has CTS, has abnormal distal motor latency. Controls: no current or past subjective complaints about upper extremities and an entirely normal neurological exam.	None reported
Rossi, 1994 ¹⁷⁸	History and symptoms typical of idiopathic CTS. Reduction of median nerve SCV in one or more of the digit-wrist segments studied, with normal values of ulnar and radial nerve sensory conduction.	Working at manual jobs. None had signs or history of cervical radiculopathy or peripheral neuropathy.
Werner, 1994 ²¹⁷	Employees at an automobile parts manufacturing plant that had reported a significant problem with CTS. Consent to testing.	Significant exposures to vibration or low temperature.
Werner, 1994 111	Referred for evaluation of CTS, must have median nerve symptoms	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Eisen, 1993 ¹⁹³	One of three groups: 1) Clinical for CTS. Symptoms and clinical signs. Examinations included Tinel's and Phalen's, but these were not required for diagnosis of CTS; 2) Historical for CTS. Symptoms: pain, sensory discomfort, or numbness in the hand, nocturnal awakening because of hand pain, clumsiness and loss of dexterity; 3) Uncertain. Vague complaints without nocturnal awakening and no loss of hand dexterity, and normal neurological exam.	1) Clinical or electrophysiological evidence of other upper limb neuropathy such as proximal median neuropathy, ulnar neuropathy, or cervical radiculopathy. 2) Historical or clinical evidence of systemic disease such as diabetes or alcoholism. 3) Prior treatment with a wrist splint or carpal tunnel surgical release. 4) Inability to obtain a median CMAP elicited by stimulating the median nerve at the wrist or inability to obtain median or ulnar SNAPs by palmar stimulation
Johnson, 1993 167	Employees at one of six poultry processing plants.	None reported
Nakamichi, 1993 ²⁶⁸	Diagnosed with carpal tunnel syndrome based on clinical signs and NCS tests. Clinical evaluation included the presence of typical sensory symptoms, Phalen's and Tinel's tests, sensory testing by 2-point discrimination on the middle finger, muscle testing, and thenar atrophy. NCS was abnormal if either DML >4.2 ms or SCV <45 m/s.	None reported
Nathan, 1993 ²¹⁸	Industrial workers from six industries: steel mill, meat/food processing, electronics, plastics, aluminum reduction, and cable plant. Workers' compensation patients had upper extremity complaints, primarily related to suspected CTS.	None reported
Rodriquez, 1993 ²⁶⁹	History and physical, and abnormal NCS	Peripheral neuropathy, cervical radiculopathy, other entrapments
Rosen, 1993 ²⁷⁰	Workers: Complaints of numbness and paresthesia and sometimes pain after long term exposure to vibrating tools. Carpal tunnel syndrome patients: Diagnosed with carpal tunnel syndrome, symptoms typical of CTS (numbness and paresthesia of radial fingers aggravated at night), not exposed to vibration	None reported
Rosén, 1993 ¹³⁸	Referred for diagnosis of suspected CTS. All had numbness and paresthesia that worsened at night	Any other explanation for symptoms, such as radiculopathy or polyneuropathy
Uncini, 1993 ¹⁶⁰	Clinical signs and symptoms suggestive of CTS, DML <4.2 ms (normal), SCV index-to-wrist >45 m/s (normal).	None reported
Buchberger, 1992 271	Patients with carpal tunnel syndrome. All had pain and sensory impairment in the distribution of the median nerve. All had prolonged DML (unspecified threshold).	None reported
Grant, 1992 ²¹⁹	Symptomatic: tingling, numbness, or decreased sensation in at least two fingers. Diagnosed: symptoms plus abnormal NCS	Arthritis, broken bones in hand/wrist, Raynaud's syndrome, previous wrist surgery, diabetes, kidney or metabolic disorders, heart or other circulatory disorders, pregnancy, use of OCs or hormones, history of heavy alcohol or tobacco use

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lmaoka, 1992 ²⁷²	Any sensory disorder in the median nerve region, and either nocturnal acroparesthesia or positive Phalen's sign.	Marked atrophy of APB, peripheral nerve disorders, diabetes, or other polyneuropathies.
Kindstrand, 1992 ²⁷³	NCS-confirmed CTS	None reported
Preston, 1992 ¹⁸⁸	Symptoms of CTS, "proven to have electrophysiologic CTS by standard nerve conduction criteria." Plus eight patients with possible CTS (symptomatic, but normal standard median studies, and at least one additional abnormal test)	None reported
Tchou, 1992 ²⁷⁴	Referred to lab with symptoms and clinically diagnosed CTS, and confirmation of diagnosis via established criteria for nerve conduction studies. Developed symptoms within three months preceding examination.	None reported
Buchberger, 1991	Symptoms of CTS.	Unrelieved or recurrent CTS after surgical treatment.
Chang, 1991 ¹⁴⁵	History of carpal tunnel syndrome, with intermittent paresthesia occurring spontaneously at night or after repetitive use of the affected hand	Diabetes
Durkan, 1991 ¹⁵⁵	Suspected carpal tunnel syndrome based on pain, numbness, and paresthesias in the distribution of the median nerve. Either abnormal motor latency or sensory latency.	None reported
Jetzer, 1991 ¹⁶⁸	One of four different groups: computer assemblers, meat processors, keyboard workers, controls.	None reported
Katz, 1991 ²⁷⁶	Pain or paresthesia in the upper extremity who were referred to the lab, and whose symptoms were caused by work.	Patients whose symptoms were not caused by work.
Lauritzen, 1991 ¹⁸⁵	Symptoms and signs compatible with CTS, and slowing of SCV along the median nerve from digit 1 or 3,or both, to the wrist, and prolonged DML from wrist to APB.	None reported
Luchetti, 1991 ¹⁶⁹	Nocturnal paresthesia in the median nerve territory. Normal motor function, sensory function, quantitative sensory examination, cutaneous trophism, distal sensory latency, distal motor latency.	Polyneuropathy, metabolic diseases with involvement of peripheral nerves.
Radwin, 1991 ¹¹⁶	Diagnosis of CTS. Sensory complaints including tingling or numbness in the thumb, index, or middle finger and nocturnal exacerbation of the paresthesias. Either positive Tinel's sign, positive Phalen's sign, or positive Semmes-Weinstein monofilaments test.	Polyneuropathy, evidence of Raynaud's phenomenon.
Charles, 1990 ¹⁷⁰	For carpal tunnel syndrome patients: Clinical diagnosis of CTS by referring physician, and at least one of the following: 1) DML = 4.5 ms; 2) median orthodromic sensory nerve conduction in the second finger <45 m/s; 3) difference between median and ulnar orthodromic distal sensory latencies in the ring finger = 0.5ms.	For controls: Diabetes, peripheral neuropathy, no symptoms suggestive of CTS For the cervical spondylitic radiculopathy group: hand paresthesia mainly in the second and third fingers
DeKrom, 1990 ²²²	Randomly selected from the general population of Maastricht (The Netherlands) and surrounding villages.	None reported

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Fitz, 1990 ²⁷⁷	APB motor latency = 4.2 ms, or digit 1 radial sensory latency = 3.1 ms, or median sensory latency = 3.2 ms or difference = 0.5 ms or similar abnormalities on digit 3	None reported
Gilliatt, 1990 278	Patients had carpal tunnel syndrome	None reported
MacDonell, 1990 ⁹⁰	Patients had at least two of five criteria: 1) DML >4.2ms; 2) SNAP amplitude <10µV; 3) SNAP conduction velocity <40m/s; 4) SNAP amplitude less than that of the ipsilateral ulnar nerve at the wrist; 5) median motor or sensory latencies at the wrist more than 0.5 ms longer than opposite hand	Normal ulnar nerve motor and sensory conduction studies in both arms
Merchut, 1990 ²⁷⁹	Symptomatic CTS referred to the lab. Electrophysiological confirmation via at least one of four NCS tests: 1) Prolonged sensory latency; 2) Prolonged DML; 3) Slowed median SCV; 4) prolonged difference between median sensory latency from ring finger and ulnar sensory latency from ring finger.	Excluded if any clinical signs, symptoms, or EMG findings suggested the possibility of another cause of paresthesia or numbness in their hands such as polyneuropathy, radiculopathy, or CNS lesion.
Palliyath, 1990 171	Symptoms of CTS, but little change on routine NCS	None reported
Pease, 1990 177	Symptoms and abnormal nerve conduction testing (vague).	Abnormalities or radial or ulnar nerves. Abnormal EMG of any muscle except the thenar muscles.
Rojviroj, 1990 ²⁸⁰	Symptoms, positive Phalen's and positive Tinel's, and carpal tunnel was confirmed by DSL >3.5 ms or DML >4.5 ms or both.	None reported
Tzeng, 1990 ¹⁸⁰	Diagnosed by both clinical and electromyographic findings	None reported
Uncini, 1990 ¹³⁵	Typical CTS symptoms but normal DML and normal or borderline SCV	None reported
Winn, 1990 ²⁸¹	Responded to ad on bulletin board	None reported
Braun, 1989 ²⁸²	Symptoms of dynamic carpal tunnel syndrome.	Evidence of long-standing fixed compression neuropathy or with contributory diseases such as rheumatoid arthrifs. Thenar atrophy or profound fixed anesthesia.
Cioni, 1989 ¹⁴⁶	Signs and symptoms suggestive of carpal tunnel syndrome. Referred to laboratory for electrophysiological confirmation of carpal tunnel syndrome.	History or physical evidence of peripheral neuropathy or cervical radiculopathy.
Jackson, 1989 ¹⁵⁰	Referred to the lab for symptoms of CTS.	Peripheral neuropathy, or obvious entrapment other than median nerve.
Meyers, 1989 ²⁸³	History and physical consistent with CTS, characteristic electrophysiologic abnormalities	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
So, 1989 ¹⁷³	Patients were selected from referrals to the lab. Carpal tunnel syndrome: Confident clinical diagnosis based on history of pain and paresthesias in the hand and fingers, and physical findings that localized the pathology to the median nerve, e.g. sensory alteration or weakness in a median nerve distribution, Tinel's, or Phalen's. Cubital tunnel syndrome: Confident clinical diagnosis based on paresthesias or numbness in an ulnar nerve distribution, usually accompanied by weakness in ulnar-innervated muscles. In those patients without weakness on examination, the diagnosis of ulnar neuropathy at the elbow was not made unless there was percussion sensitivity at the cubital tunnel or the ulnar groove, or exacerbation of symptoms with elbow flexion.	None reported
Szabo, 1989 ²⁸⁴	CTS patients about to have carpal tunnel release surgery. Clinical and electrophysiological evidence of CTS. Electrophysiological evidence based on either DML >4.5 ms or DSL >3.5 ms.	None reported
Uncini, 1989 ¹⁶¹	Symptoms and signs of carpal tunnel syndrome	Severe carpal tunnel (DML >4.2 ms or SNAPs were absent or SNAPs were very low amplitude)
De Léan, 1988 ²⁸⁵	Paresthesia in median nerve distribution, regardless of Tinel's or Phalen's signs	Polyneuropathy, medicolegal cases, workers' comp
Koris, 1988 ¹⁹⁸	Accepted signs and symptoms including paresthesia, but did not have to be limited to the median nerve distribution	None reported
Molitor, 1988 110	Referred to lab for the diagnosis of carpal tunnel.	None reported
Mortier, 1988 ²⁸⁶	Prolonged distal mobr latency of median nerve or prolonged distal sensory latency of median nerve	Generalized peripheral neuropathy, other peripheral entrapment neuropathies, cervical radiculopathy.
Pease, 1988 ²⁸⁷	Diagnosed with CTS based on clinical and electrodiagnostic findings	None reported
Carroll, 1987 ²⁸⁸	Referred to lab, symptoms suggestive of CTS	Abnormal ulnar sensory amplitude or latency.
Jessurun, 1987 ²⁸⁹	Suffering from primary CTS	None reported
Johnson, 1987 ²⁹⁰	Antidromic DSL to middle finger >4 ms and DML >4.3 ms.	None reported
Liang, 1987 ²⁹¹	None reported	None reported
Macleod, 1987 ²⁹²	Symptomatic NCS confirmed with abnormal sensory latency	Signs of other neurologic disorder
Seror, 1987 156	Pathological wrists	Radicular signs
Borg, 1986 ²⁹³	Referred to lab with suspicion of CTS. Patients had digital paresthesias.	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Gellman, 1986 ¹⁰⁶	Carpal tunnel group syndrome: Three requirements: 1) Symptoms indicative of median-nerve compression in the carpal canal; 2) Either positive Semmes-Weinstein test or positive two-point discrimination test; 3) Positive nerve conduction results as indicated by any of four abnormalities: A) DML >4.5 ms B) DML on symptomatic hand more than 1 ms slower than DML on asymptomatic hand C) Sensory latency >3.5 ms D) Sensory latency on symptomatic hand more than one millisecond slower than on asymptomatic hand. Diverse lesion group: Abnormal results on clinical sensibility testing other than carpal tunnel syndrome	None reported
Escobar, 1985 ¹⁵¹	Patients: Referred to lab for evaluation of numbness, tingling, weakness, and/or pain in the hand or arm. Controls: DSL <3.7 ms.	Endocrine disorders or peripheral nerve disease.
Kimura, 1985 ¹⁸⁹	Referred to lab with frank clinical signs and symptoms suggestive of CTS	Other disease that predispose toward peripheral neuropathy.
Mills, 1985 ¹⁹⁴	Tentative diagnosis of CTS	None reported
Borg, 1984 ²⁹⁴	Patients with CTS. Some patients' conditions had	None reported
Pryse-Phillips, 1984	been neurophysiologically confirmed (undefined). Group 01: Carpal tunnel syndrome: Symptoms of paresthesia, numbness and/or weakness in the hand in digits I-II or I-V, with or without hand and arm pain, usually with nocturnal or early morning accentuation, ± clinical signs of thenar motor or median nerve territory sensory deficit. DML >4.5 ms or a difference of 1 ms between right and left or 1.5 median/ulnar difference. Median SNAP amplitude <ulnar <10="" latency="" onset="" or="" to="" µv="">3.5 ms. Group 02: Cubital tunnel syndrome: Symptoms of hand weakness, ± digit V (IV) hypoesthesia, not extending into palm: and/or electrical signs of interosseous or hypothenar wasting, with proportionate weakness. Eisen score (undefined) greater than 5/10. Group 03: Other median nerve pathologies: Digital neuropathy affecting digits I-III or arm pain/paresthesia without nocturnal predominance, or clinically apparent weakness of long forearm flexors, ± palmar hypoesthesia. EMG evidence of acute/chronic denervation in forearm flexor muscles, ± delay in motor conduction across the point above the wrist with absence of electrical evidence of median nerve compression at the carpal tunnel. Group 04: Thoracic outlet syndrome. Group 05: Cervical radiculopathy</ulnar>	Carpal tunnel syndrome: Martin-Gruber anastomosis, other median nerve pathologies: cases of anterior interosseous syndrome
Satoh, 1984 ²⁹⁵	No symptoms, normal ulnar sensory and motor conduction and one of three nerve conduction abnormalities: 1) orthodromic SCV digit to-palm <42 m/s; 2) terminal latency >4.2 ms; 3) absent SNAP and absent CMAP.	None reported

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Szabo, 1984 ³⁰	Patients with objectively proved abnormalities of median nerve conduction who had carpal tunnel release surgery.	None reported
Goddard, 1983 ²⁹⁶	Diagnosed with CTS and referred to the department	None reported
Kim, 1983 ¹⁹⁵	Signs and symptoms highly suggestive of CTS but with borderline or normal DSL.	None reported
Marin, 1983 ¹³⁹	Patients had previously undergone routine NCS studies for carpal tunnel syndrome	None reported
Wongsam, 1983 172	Symptoms suggesting early CTS	None reported
Johnson, 1981 ²⁹⁷	Diagnosed CTS: history and NCS	None reported
Dekel, 1980 ²¹	Diagnosed with carpal tunnel using history, clinical exam, and nerve conduction studies.	Any of the recognized diseases associated with carpal tunnel syndrome.
Messina, 1980 120	Signs and symptoms suggestive of CTS	None reported
Gelmers, 1979 ²⁹	Diagnosis of carpal tunnel based on three findings: 1) Acroparesthesia in the distribution of the median nerve; 2) Thenar muscle wasting or weakness or failure to detect an action potential of the thenar muscles by needle electromyography; 3) Prolongation of distal latency of the median nerve to more than 4.7 ms, or a difference in distal latency of more than 1 ms between symptomatic and asymptomatic hands, even though both latencies were within normal limits	Signs of generalized neuropathy
Kimura, 1979 ¹⁴⁰	Clinical impression (history and symptoms, not NCS), relatively mild symptoms	Polyneuropathy
Schwartz, 1979 ¹⁸⁷	Referred to lab based on sensory symptoms in a median distribution.	Generalized neuropathy
Stewart, 1978 157	In addition to ipsilateral ulnar sensory amplitude = $8.5 \mu V$ and ulnar sensory latency < $2.8 \mu V$ ms, three or more of the following were required: 1) Sensory signs in the distribution of the median nerve.; 2) Thenar wasting or weakness; 3) DML > $4.5 \mu V$ ms; 4) sensory onset latency > $2.7 \mu V$ sensory amplitude < $8.6 \mu V$	Diabetes, peripheral neuropathy. CTS secondary to trauma or other localized or generalized disease.

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Eisen, 1977 ²⁹⁸	Carpal tunnel patients: Sensory symptoms limited to one or both hands, normal ulnar sensory latency (<2.8 ms), normal ulnar sensory amplitude (>8.4 µV), and at least three of the following five criteria: 1) Sensory signs restricted to median distribution; 2) Weakness or wasting of the APB muscle; 3) Median DML >4.5 ms; 4) Median DSL >2.7 ms; 5) Median SNAP amplitude <8.6 µV or median SNAP duration >2.4 ms. Cubital tunnel patients: Sensory symptoms limited to one or both hands, normal median sensory latency (<2.7 ms), normal median sensory amplitude (>8.6 µV), and at least three of the following six criteria: 1) Sensory signs restricted to ulnar distribution; 2) Weakness or wasting of the ulnar-innervated muscles of the hand; 3) Ulnar DML >4.0 ms; 4) Ulnar proximal motor latency (stimulation just above the elbow) >8.9 ms; 5) Ulnar DSL >2.8 ms; 6) Ulnar SNAP amplitude <8.4 µV or ulnar SNAP duration >2.1 ms. Patients with proximal lesions: Sensory symptoms limited to one or both hands, but did not meet criteria for either carpal tunnel or cubital tunnel.	Subjects were excluded from the control group if there was neuromuscular disease, diabetes, alcoholism, peripheral neuropathy, or systemic dysfunction.
Sedal, 1973 ²⁹⁹	Presented as idiopathic carpal tunnel.	Excluded if CTS was an incidental finding in the investigation of a generalized peripheral neuropathy, OR if they had diabetes or alcoholism or chronic renal disease, or if there was clinical evidence of either radial or nerve lesions
Welch, 1973 ²²³	Workers at a factory employed on repetition work producing domestic appliances. The other group consisted of job applicants who had not yet started work.	None reported
Casey, 1972 300	Carpal tunnel syndrome: Classical symptoms. Also 10 of the 16 patients had hypalgesia in the fingers of the involved hand supplied by the median nerve. Abnormal (or at the lower limit of normal) median SNAP recorded at the wrist after digital stimulation. Diabetics: Reflex changes and distal sensory abnormalities in the lower limbs, consisting of pain and paresthesia with sensory loss. In addition, 10 of the 18 diabetics had sensory changes in the upper limbs	None reported
Loong, 1972 141	Clinical diagnosis of CTS with typical history of intermittent paresthesia at night or after use.	None reported
Melvin, 1972 ¹⁴⁷	Referred to the laboratory as possible cases of carpal tunnel syndrome.	None reported
Buchthal, 1971 ³⁰¹	None reported	Normal ulnar SCV and latency to ADM to exclude generalized neuropathy

Article	Reported Patient Inclusion Criteria	Reported Patient Exclusion Criteria
Loong, 1971 ¹⁴⁸	Referred to lab with clinical diagnosis of carpal tunnel syndrome. Typical history of the syndrome with intermittent paresthesia occurring spontaneously at night or after use of the affected hand.	Diabetes
Plaja, 1971 ¹⁴²	None reported	"We excluded misleading diagnosis by controlling at the same time different levels and nerve trunks."

Question #2: What are the specific indications for surgery for carpal tunnel syndrome?

Published evidence does not directly address the specific indications for surgery for carpal tunnel syndrome. Therefore, we describe the reported characteristics of patients who have received surgery for carpal tunnel syndrome in published studies. The extent to which these patients represent typical surgical candidates is not certain. Patients included in published studies of a procedure are frequently a subset of patients who are candidates for that procedure. They may represent an unusual group of interest, or a group thought most likely to benefit from the procedure. Therefore, the data presented here, while informative, may not accurately reflect the overall patient population. It does, however, represent the best data available, and is the most comprehensive description of those carpal tunnel syndrome patient characteristics who receive surgery that has yet been compiled.

Evidence Base

To answer this question, we examined 141 studies (controlled trials and case series) describing a total of 15,993 patients.

Age

Patients who received surgery for carpal tunnel syndrome were predominantly of middle age. The mean of mean ages from the 124 studies that reported this information was 50.5 years, with a standard deviation of 5.7. Ages of individual patients ranged from 17 to 100 years. Mean ages and ranges from individual studies are given in Table 47, and are depicted in Figure 16. The vertical line in Figure 16 represents the mean age for all studies.

Very few studies (4%) reported that patients were excluded on the basis of age. Two studies excluded patients under the age of 18, 302,303 and one excluded patients under $16.^{304}$ In contrast, one excluded patients over the age of seventy, 305 and another excluded patients over $75.^{306}$

Sex

Patients receiving surgery for carpal tunnel syndrome were more likely to be female than male, as can be seen in Figure 17. One hundred twenty eight studies provided sufficient information to calculate the male-to-female patient ratio. The average study reported that 73% of patients were female, with a standard deviation of 0.2. Patients in two studies were 100% female, and 100% male in one study. Numbers of male and female patients in individual studies are reported in Table 47.

No study reported sex to be a criterion for exclusion or inclusion. However, both studies in which men were the majority recruited their patients from male-majority populations. One recruited exclusively from a veteran's hospital population, ³⁰⁷ and one

recruited patients who worked with heavy, vibrating machinery.³⁰⁸ These patients do not represent typical carpal tunnel syndrome patients.

Signs and Symptoms

Signs and symptoms of carpal tunnel syndrome among patients receiving surgery for carpal tunnel syndrome were incompletely reported. This is illustrated by Figure 18, which depicts the percentage of studies reporting the number of patients with an individual sign or symptom. This percentage never exceeds 15% of all studies. Rather than report the number of patients with a given sign or symptom, the common practice among studies of carpal tunnel syndrome is to report that patients had one or more symptoms from a given list. Some studies that included patients with bilateral carpal tunnel syndrome report symptoms per affected hand rather than per patient, reflecting the fact that the same patient can have different symptoms in each hand. The number and percent of patients reporting each sign or symptom is given in Table 48. These data are summarized in Figure 19. "Error" bars in Figure 19 represent the range of percentages reported by individual studies. Because so few (always less than 15%) studies reported this information, the extent to which the available data reflect the signs and symptoms of typical patients receiving surgery cannot be determined.

Eight studies excluded patients with thenar atrophy, while four included only patients with thenar atrophy. Seven studies required their patients to have Tinel's sign, Phalen's sign or both, and an indeterminate number included tests for these signs as part of their diagnostic procedure. The exact number of such studies can not be determined because some describe their patients as having "signs and symptoms of carpal tunnel syndrome" without providing further description or enumeration. The extent to which use of these criteria influence the overall description of the typical patient with carpal tunnel syndrome cannot be determined, because it is unclear whether or to what extent criteria for surgery may differ from criteria for study inclusion.

The duration of symptoms prior to surgery was reported by 35 studies (24% of total). These are listed in Table 49. The mean of means among these 35 studies was 29.9 months, with a standard deviation of 16.5 and a range of zero to 480 months. The means and ranges of individual studies are depicted in Figure 20. The vertical line in Figure 20 represents the mean of means.

Neuroelectrical characteristics

Of the 145 studies that reported on surgery for carpal tunnel syndrome and met inclusion criteria, 83 stated that electrodiagnostic tests were part of their inclusion criteria, but did not provide any further information as to the nature of these tests. An additional 26 did not provide any diagnostic information. Eleven studies did not include electrodiagnostic studies in their description of their diagnostic and inclusion criteria, and two specifically stated that electrodiagnostics were not part of their diagnostic protocol. Electrodiagnostic criteria in the remaining studies are reported in Table 50. Because the majority of studies excluded some patients based on their

Table 47. Age and sex of patients receiving surgery for carpal tunnel syndrome

Trial	Number of patients	Number of males	Number of females	Percent female	Age reported as mean or median?	Age	Age of youngest patient	Age of oldest patient
Finsen, 2001	79	18	61	77.2%	Median	48	21	86
Mondelli, 2001	28	4	24	85.7%	Mean	52.8	35	75
Avci, 2000 309	25	1	24	96.0%	Mean	43	21	72
Khan, 2000 310	44	11	33	75.0%	Mean	55	29	88
Mondelli, 2000 311	110	13	97	88.2%	Mean	56	20	82
Muller, 2000 312	148	28	120	81.1%	Mean	51.8	NRa	NR
Porras, 2000	85	8	77	90.6%	Mean	52	18	81
Vartimidis, 2000 314	15	6	9	60.0%	Mean	52	28	75
Alderson, 1999	26	5	21	80.8%	Mean	44.4	22	79
Braun, 1999 ³¹⁶	225	36	189	84.0%	Mean	41.0	NR	NR
Chen, 1999 ³¹⁷	948	212	736	77.6%	Mean	48	21	79
Erhard, 1999	124	15	109	87.9%	Mean	54.3	19	84
Finsen, 1999	82	22	60	73.2%	Mean	49.4	21	86
Hasegawa, 1999 ³²⁰	82	0	82	100.0	Mean	54.1	NR	NR
Hirooka, 1999	37	4	33	89.2%	Mean	58	40	78
Lindau, 1999 322	140	17	123	87.9%	Mean	55.4	NR	NR
Olney, 1999 323	211	46	165	78.2%	Mean	44.8	NR	NR
Senda, 1999 324	26	1	25	96.2%	Mean	56.8	19	93
Straub, 1999 305	67	47	20	29.9%	Median	40	19	70
Vartimidis, 1999 325	22	8	14	63.6%	Mean	52	21	77
Atroshi, 1998 326	103	35	68	66.0%	Mean	52	21	88
Aulisa, 1998 327	45	8	37	82.2%	Mean	47	26	68
Buckhorn 1998	50	21	29	58.0%	Mean	51.3	27	61
Choi, 1998 329	154	6	148	96.1%	Mean	52	30	82
Davies, 1998 330	239	NR	NR	NR	Mean	43.5	20	82
Lee, 1998 331	525	134	391	74.5%	Mean	50.7	21	88
Nakamichi, 1998 ³³²	130	16	114	87.7%	Mean	58	35	85

Trial	Number of	Number of males			Age reported	Age	youngest	Age of oldest
	patients		of females	Percent female	as mean or median?		patient	patient
Papageorgiou, 1998 333	76	18	58	76.3%	Mean	48	NR	NR
Schuind. 1998 334	13	6	7	53.8%	Mean	47	45	77
Tomaino, 1998 335	29	6	23	79.3%	Mean	52	28	82
Armstrong, 1997 ³³⁶	176	35	141	80.1%	Mean	50.5	30	86
Atroshi, 1997 337	204	56	148	72.5%	Mean	49.3	19	94
Baguneid, 1997 338	75	11	64	85.3%	Mean	56	24	85
Chia, 1997 339	62	13	49	79.0%	Mean	47.7	29	73
Citron, 1997 340	47	8	39	83.0%	Mean	52.1	26	80
Higgs, 1997 341	93	30	63	67.7%	Mean	43	23	69
Karlsson, 1997	74	15	59	79.7%	Median	54.5	24	88
Katz, 1997 302	135	42	93	68.9%	NR	NR	NR	NR
Leinberry, 1997 ³⁴²	44	18	26	59.1%	Mean	64.9	38	100
Rosen, 1997	102	18	84	82.4%	Mean	51.0	24	82
Serra, 1997 344	112	16	96	85.7%	Mean	47	31	70
Stahl, 1997 345	50	16	34	68.0%	Mean	49.5	NR	NR
Tucci, 1997 346	27	6	21	77.8%	Mean	48.6	NR	NR
Weber, 1997 347	74	26	48	64.9%	Median	41.4	26	80
Wheatly, 1997	126	114	12	9.5%	NR	NR	NR	NR
Cobb, 1996 348	235	44	191	81.3%	Mean	51	20	79
Elmaraghy. 1996 ³⁴⁹	69	21	48	69.6%	Mean	51	24	97
Franzini, 1996 350	50	11	39	78.0%	Mean	52	32	60
Gibbs, 1996 351	46	16	30	65.2%	Mean	56.2	31	86
Glowacki, 1996 352	167	35	132	79.0%	Mean	42	17	84
Jacobsen, 1996 353	32	9	23	71.9%	Mean	44.9	24	59
Kluge. 1996 354	66	18	48	72.7%	Mean	51	36	93
Lee, 1996 355	275	76	199	72.4%	Mean	50.7	21	88
Mclaughlin, 1996 ³⁵⁶	102	26	76	74.5%	Mean	52	NR	NR
Nagle, 1996 357	506	134	372	73.5%	Mean	48	13	91
Nygaard, 1996 306	29	7	22	75.9%	Mean	53	32	75
Okutsu, 1996 41	43	2	41	95.3%	Mean	55.1	31	87

Trial	Number of patients	Number of males	Number of females	Percent female	Age reported as mean or median?	Age	Age of youngest patient	Age of oldest patient
Padua, 1996 358	33	7	26	78.8%	Mean	47.2	NR	NR
Pennino, 1996 359	124	NR	NR	NR	Mean	55	28	92
Povlsen, 1996 360	51	23	28	54.9%	NR	NR	NR	NR
Strickland, 1996 ³⁶¹	62	16	46	74.2%	Mean	52	22	88
Wintman, 1996 362	50	NR	NR	NR	Mean	54	25	83
Worseg, 1996	126	38	88	69.8%	Mean	56.0	35	90
Abdullah, 1995 363	100	19	81	81.0%	Mean	41.4	19	79
Bury, 1995 364	43	4	39	90.7%	Mean	52.3	NR	NR
Dumontier, 1995 ³⁶⁵	96	11	85	88.5%	Mean	41.1	29	53
El-Zahaar, 1995 ⁴³	41	12	29	70.7%	Mean	53	39	61
Futami, 1995 366	10	1	9	90.0%	Mean	51	NR	NR
Gross, 1995 367	44	16	28	63.6%	Mean	44.2	NR	NR
Hallock, 1995 368	100	26	74	74.0%	Mean	59	NR	NR
Katz, 1995 369	50	6	44	88.0%	Mean	51.4	NR	NR
Lang, 1995 109	23	5	18	78.3%	Mean	53	25	84
LoVerme, 1995 370	42	4	38	90.5%	Mean	29	NR	NR
Mirza, 1995 ³⁷¹	236	74	162	68.6%	Mean	44	17	79
Nancollas, 1995 ³⁷²	93	17	76	81.7%	Mean	52.5	NR	NR
Sennwald, 1995 ³⁷³	47	12	35	74.5%	Mean	54	22	88
Shinya, 1995 374	88	16	72	81.8%	Mean	49	20	82
Al-Qattan, 1994 375	112	28	84	75.0%	Mean	54	25	83
Chow, 1994 42	815	289	526	64.5%	NR	NR	NR	NR
Erdmann, 1994 304	96	26	70	72.9%	Mean	53.4	NR	NR
Foulkes, 1994 376	33	16	17	51.5%	Mean	45.4	NR	NR
Katz, 1994 377	104	31	73	70.2%	Mean	55	25	87
Kelly, 1994 378	69	16	53	76.8%	Mean	50	21	79
Kerr, 1994 379	85	37	48	56.5%	Mean	44.8	19	82
Menon, 1994 380	87	28	59	67.8%	Mean	48.3	21	76

Trial	Number of	Number of males			Age reported	Age	youngest	Age of oldest
	patients		of females	Percent female	as mean or median?		patient	patient
Pascoe, 1994 381	28	12	16	57.1%	Mean	55	32	82
Payne, 1994 382	16	6	10	62.5%	NR	NR	NR	NR
Roth. 1994 383	94	35	59	62.8%	Mean	52.4	25	91
Singh, 1994 384	357	56	301	84.3%	NR	NR	NR	NR
Skoff, 1994 385	1994	NR	NR	NR	Mean	56.0	24	84
Slattery, 1994 40	215	69	146	67.9%	Mean	41	17	84
Strasberg, 1994 ³⁸⁶	45	16	29	64.4%	Mean	50.6	NR	NR
Wolson, 1994 387	30	10	20	66.7%	Mean	47	14	71
Biyani, 1993 ³⁸⁸	56	7	49	87.5%	Mean	65.4	44	81
Brown, 1993 ⁴⁵	145	46	99	68.3%	Mean	55	25	87
Chang, 1993	30	6	24	80.0%	Mean	46.2	31	77
Feinstein, 1993	55	21	34	61.8%	Mean	45	21	79
Jiminez, 1993 391	24	6	18	75.0%	Mean	46	NR	NR
Leach, 1993 392	25	11	14	56.0%	Mean	43	25	80
Levine, 1993	39	17	22	56.4%	Median	57	19	88
Nakamichi, 1993 ³⁹⁴	41	8	33	80.5%	Mean	54	33	86
Nathan, 1993 395	238	80	158	66.4%	Mean	41	15	79
Okutsu, 1993 396	27	0	27	100.0%	Mean	55.9	33	87
Palmer, 1993	173	73	100	57.8%	Mean	44.9	20	83
Waegeneers, 1993 ³⁹⁸	76	21	55	72.4%	Mean	54	21	82
Nolan, 1992 399	22	7	15	68.2%	Mean	70	52	86
Pagnanelli, 1992 400	228	65	163	71.5%	Mean	55.2	NR	NR
Viegas, 1992 401	71	17	54	76.1%	Mean	48	23	79
Young, 1992	21	NR	NR	NR	Mean	49	22	72
Yu, 1992 ⁴⁰³	53	22	31	58.5%	Median	46	20	83
Flaschka, 1991	99	18	81	81.8%	Mean	56.4	22	82
Foucher, 1991	83	17	66	79.5%	Mean	59.6	46	77
Hagberg, 1991	41	41	0	0.0%	Mean	42.0	NR	NR

Trial	Number of patients	Number of males	Number of females	Percent female	Age reported as mean or median?		youngest patient	Age of oldest patient
Jakab, 1991 406	73	25	48	65.8%	Mean	52	27	88
Mackimmon, 1991 ⁴⁰⁷	59	11	48	81.4%	Mean	58.5	20	91
Resnick, 1991 408	65	17	48	73.8%	Mean	46.2	23	81
Schuind, 1990 409	21	2	19	90.5%	Mean	49	32	81
Gellman, 1989 410	21	2	19	90.5%	Mean	51.5	30	65
Okutsu, 1989	45	15	30	66.7%	Mean	51.1	29	73
Richman, 1989	12	6	6	50.0%	NR	NR	NR	NR
Seiler, 1989 413	10	2	8	80.0%	Mean	43.6	23	65
Seradge, 1989	500	218	282	56.4%	Median	41	19	87
Szabo, 1989 284	22	6	16	72.7%	Mean	51	24	79
Gelberman, 1987 415	29	17	12	41.4%	Mean	55	28	84
Holmgren, 1987 416	48	15	33	68.8%	Mean	50	21	80
Gartsman, 1986 417	50	14	36	72.0%	NR	NR	NR	NR
Kulick, 1986 418	167	30	137	82.0%	Mean	55.5	21	92
Leblhuber , 1986 419	47	10	37	78.7%	Mean	50.2	19	81
Shurr, 1986 420	36	8	28	77.8%	Mean	44.6	NR	NR
Wadstroem, 1986 ⁴²¹	36	10	26	72.2%	Mean	50	32	80
Rhodes, 1985	32	21	11	34.4%	Mean	63	37	90
Litchman, 1984	135	28	107	79.3%	Mean	54	20	84
van Rossum, 1980 ⁴²⁴	37	6	31	83.8%	NR	NR	NR	NR

a: Not reported

Table 48. Symptoms of patients treated with surgery for carpal tunnel syndrome

Study	Number of patients (or	Sign or symptom Number of patients with sign or		Percent of patients (or hands)
Mol quablin	hands)	Durning	symptom 70	40.40/
McLaughlin, 1996 356	102	Burning	70	68.6%
Mirza, 1995 ³⁷¹	56	Burning	6	10.7%
Finsen, 2001	79	Clumsiness	42	53.2%
224	, ,	Cidinisinoss		00.270
Atroshi, 1997	255 Hands	Clumsiness	155	60.8%
Cobb, 1996 348	235	Clumsiness	81	34.5%
Lee, 1996 355	275 Hands	Clumsiness	207	75.3%
Lascar, 2000 425	71	Clumsiness	6	8.5%
Porras, 2000	85	Durkan/carpal compression test	50	58.8%
Finsen, 2001	79	Night symptoms	56	70.9%
Straub, 1999	100 Hands	Night symptoms	93	93.0%
Aulisa, 1998 327	45	Night symptoms	44	97.8%
Buchhorn, 1998 ³²⁸	50	Night symptoms	50	100.0%
Atroshi, 1997	255 Hands	Night symptoms	237	92.9%
Cobb, 1996 348	235	Night symptoms	71	30.2%
Elmaraghy, 1996 349	69	Night symptoms	56	81.2%
Glowacki, 1996				
352	167	Night symptoms	114	68.3%
Kluge, 1996 354	66	Night symptoms	50	75.8%
Lee, 1996 355	275 Hands	Night symptoms	226	82.2%
McLaughlin, 1996 ³⁵⁶	102	Night symptoms	78	76.5%
Nygaard, 1996 306	29	Night symptoms	20	69.0%
Strickland, 1996 361	58	Night symptoms	58	100%
Worseg, 1996				
44	126	Night symptoms	111	88.1%
Singh, 1994 384	357	Night symptoms	104	29.1%
Palmer, 1993	173	Night symptoms	148	85.5%
Pagnanelli, 1992 400	456 Hands	Night symptoms	424	93.0%
Resnick, 1991	75 Hands	Night symptoms	66	88.0%

Study	of symptom patients patients (or or		with sign	Percent of patients (or hands)
Freshwater, 1978 426	22	Night symptoms	22	100%
Provinciali, 2000 427	100	Numbness	62	62.0%
Vartimidis, 2000 314	15	Numbness	15	100.0%
Straub, 1999 305	100 Hands	Numbness	71	71.0%
Aulisa, 1998 327	45	Numbness	7	15.6%
Armstrong, 1997 ³³⁶	208 Hands	Numbness	160	76.9%
Atroshi, 1997	255 Hands	Numbness	178	69.8%
Blair, 1996 428	75	Numbness	71	94.7%
Cobb, 1996 348	235	Numbness	88	37.4%
Elmarghy, 1996	69	Numbness	68	98.6%
Kluge, 1996 354	66	Numbness	35	53.0%
Lee, 1996 355	275 Hands	Numbness	240	87.3%
McLaughlin, 1996 356	102	Numbness	71	69.6%
Futami, 1995	10	Numbness	10	100%
LoVerme, 1995	42	Numbness	28	66.7%
Mirza, 1995 371	56	Numbness	53	94.6%
Singh, 1994 384	357	Numbness	283	79.3%
Strasberg, 1994 386	45	Numbness	45	100.0%
Waegeneers, 1993 398	100 Hands	Numbness	28	28.0%
Pagnanelli, 1992 400	456 Hands	Numbness	264	57.9%
Wadstroem, 1986 421	36	Numbness	25	69.4%
Freshwater, 1978 426	11	Numbness	11	100%
Provinciali, 2000 ⁴²⁷	100	Pain	80	80.0%
Vartimidis, 2000 314	15	Pain	15	100%
Armstrong, 1997 336	208 Hands	Pain	185	88.9%
Atroshi, 1997	255 Hands	Pain	198	77.6%
Blair, 1996 428	75	Pain	67	89.3%
Cobb, 1996 348	131	Pain	80	61.1%

Study	Number of patients (or	Sign or symptom	Number of patients with sign or	Percent of patients (or hands)
F	hands)		symptom	05 50/
Elmaraghy, 1996 349	69	Pain	59	85.5%
Lee, 1996 355	275 Hands	Pain	232	84.4%
Mirza, 1995 371	56	Pain	46	82.1%
Strasberg, 1994 ³⁸⁶	45	Pain	39	86.7%
Waegeneers. 1993 398	100 Hands	Pain	96	96.0%
Nolan, 1992 399	22	Pain	11	50.0%
Richman, 1989	12	Pain	10	83.3%
Lowry, 1988 429	50	Pain	47	94.0%
Freshwater, 1978 426	22	Pain	6	27.3%
Nygaard, 1996 306	29	Paresis	8	27.6%
Provinciali, 2000 427	100	Paresthesias	82	82.0%
Straub, 1999 305	100 Hands	Paresthesias	100	100%
Buchholm, 1998 328	50	Paresthesias	49	98.0%
Armstrong, 1997 ³³⁶	208 Hands	Paresthesias	195	93.8%
Atroshi, 1997	255 Hands	Paresthesias	242	94.9%
Cobb, 1996 348	235	Paresthesias	82	34.9%
Elmaraghy, 1996 349	69	Paresthesias	59	85.5%
Kluge, 1996 354	66	Paresthesias	3	4.5%
Lee, 1996 355	275 Hands	Paresthesias	233	84.7%
Worseg, 1996	126	Paresthesias	120	95.2%
Mirza, 1995 371	56	Paresthesias	56	100%
Palmer, 1993	173	Paresthesias	171	98.8%
Waegeneers, 1993 398	100 Hands	Paresthesias	99	99.0%
Pagnanelli, 1992 400	456 Hands	Paresthesias	424	93.0%
Wadstroem, 1986 421	36	Paresthesias	32	88.9%
Finsen, 2001	79	Phalen's sign	58	73.4%
Porras, 2000	85	Phalen's sign	64	75.3%

Study	Number of patients (or hands)	Sign or symptom	Number of patients with sign or symptom	Percent of patients (or hands)
Straub, 1999 305	100 Hands	Phalen's sign	87	87.0%
Aulisa, 1998 327	45	Phalen's sign	32	71.1%
Atroshi, 1997	255 Hands	Phalen's sign	214	83.9%
Serra, 1997 344	112	Phalen's sign	98	87.5%
Glowacki, 1996 352	167	Phalen's sign	115	68.9%
McLaughlin, 1996 356	102	Phalen's sign	90	88.2%
Nygaard, 1996 306	29	Phalen's sign	22	75.9%
Strickland, 1996 361	62	Phalen's sign	45	72.6%
Worseg, 1996				
44	126	Phalen's sign	74	58.7%
Bury, 1995 364	43	Phalen's sign	43	100.0%
Futami, 1995	10	Phalen's sign	10	100.0%
Lang, 1995 109	23	Phalen's sign	19	82.6%
Erdmann. 1994 304	96	Phalen's sign	80	83.3%
Payne, 1994 382	16	Phalen's sign	16	100.0%
Roth, 1994 383	94	Phalen's sign	94	100.0%
Palmer, 1993				
397	211 Hands	Phalen's sign	196	92.9%
Waegemeers, 1993 398	100 Hands	Phalen's sign	84	84.0%
Resnick, 1991	75 Hands	Phalen's sign	69	92.0%
Richman, 1989	12	Phalen's sign	10	83.3%
Freshwater, 1978 426	22	Phalen's sign	17	77.3%
Armstrong, 1997 ³³⁶	208 Hands	Stiffness	174	83.7%
Lascar, 2000 425	71	Stiffness	7	9.9%
Aulisa, 1998 327	45	Swelling	27	60.0%
Mirza, 1995 371	280	Swelling	3	1.1%
Freshwater, 1978 426	22	Swelling	0	0.0%
Strickland, 1996 361	58	Tenderness	54	93.1%
Pagnanelli, 1992 400	456 Hands	Tenderness	18	3.9%

Study	Number of patients (or hands)	Sign or symptom	Number of patients with sign or symptom	Percent of patients (or hands)
Porras, 2000 313	85	Thenar atrophy	15	17.6%
Aulisa, 1998 327	45	Thenar atrophy	3	6.7%
Buchhorn, 1998 ³²⁸	50	Thenar atrophy	11	22.0%
Atroshi, 1997	255 Hands	Thenar atrophy	36	14.1%
Serra, 1997 344	112	Thenar atrophy	16	14.3%
McLaughlin, 1996 ³⁵⁶	102	Thenar atrophy	16	15.7%
Nygaard, 1996 306	29	Thenar atrophy	8	27.6%
LoVerme, 1995	42	Thenar atrophy	8	19.0%
Singh, 1994 384	357	Thenar atrophy	110	30.8%
Waegeneers, 1993 398	100 Hands	Thenar atrophy	8	8.0%
Nolan, 1992 399	22	Thenar atrophy	11	50.0%
Pagnanelli, 1992 400	456 Hands	Thenar atrophy	112	24.6%
Foucher, 1991 405	83	Thenar atrophy	83	100.0%
Mackimmon,				40.504
1991 ⁴⁰⁷	59	Thenar atrophy	41	69.5%
Resnick, 1991	75 Hands	Thenar atrophy	12	16.0%
Richman, 1989	12	Thenar atrophy	3	25.0%
Gelberman, 1987 415	61	Thenar atrophy	38	62.3%
Kulick, 1986 418		Thenar atrophy	20	12.0%
Leblhuber, 1986 419	55 Hands	Thenar atrophy	14	25.5%
Wadstroem, 1986 421	36	Thenar atrophy	17	47.2%
Freshwater, 1978 426	22	Thenar atrophy	2	9.1%
Finsen, 2001	79	Tinel's sign	46	58.2%
Porras, 2000	85	Tinel's sign	51	60.0%
Straub, 1999 305	100 Hands	Tinel's sign	73	73.0%
Buchhorn, 1998 ³²⁸	50	Tinel's sign	46	92.0%
Atroshi, 1997	255 Hands	Tinel's sign	176	69.0%

Study	Number of patients (or	Sign or symptom	Number of patients with sign or	Percent of patients (or hands)
	hands)		symptom	
Serra, 1997 344	112	Tinel's sign	5	4.5%
Glowacki, 1996				
352	96	Tinel's sign	66	68.8%
McLaughlin, 1996 356	102	Tinel's sign	69	67.6%
Nygaard, 1996 306	29	Tinel's sign	9	31.0%
Strickland, 1996 361	62	Tinel's sign	45	72.6%
Worsegm 1996				
44	126	Tinel's sign	100	79.4%
Futami, 1995	10	Tinel's sign	10	100.0%
Lang, 1995 109	23	Tinel's sign	7	30.4%
Erdmann, 1994				
304	96	Tinel's sign	74	77.1%
Roth, 1994 383	94	Tinel's sign	94	100.0%
Palmer, 1993	211	Tinel's sign	181	85.8%
Waegeneers, 1993 ³⁹⁸	100 Hands	Tinel's sign	77	77.0%
Resnick, 1991	75 Hands	Tinel's sign	57	76.0%
Richman, 1989	12	Tinel's sign	7	58.3%
Freshwater,				
1978 426	22	Tinel's sign	15	68.2%
Provinciali, 2000 ⁴²⁷	100	Weakness	75	75.0%
Straub, 1999 305	100 Hands	Weakness	63	63.0%
Aulisa, 1998 327	45	Weakness	9	20.0%
Armstrong, 1997 ³³⁶	208 Hands	Weakness	156	75.0%
Atroshi, 1997	255 Hands	Weakness	79	31.0%
Cobb, 1996 348	235	Weakness	97	41.3%
Elmaraghy, 1996 349	69	Weakness	35	50.7%
Kluge, 1996 354	66	Weakness	5	7.6%
Lee, 1996 355	275 Hands	Weakness	220	80.0%
McLaughlin, 1996 ³⁵⁶	102	Weakness	17	16.7%
Singh, 1994 384	357	Weakness	120	33.6%
Strasberg, 1994 ³⁸⁶	45	Weakness	42	93.3%

Study	Number of patients (or hands)	Sign or symptom	Number of patients with sign or symptom	Percent of patients (or hands)
Palmer, 1993				
397	173	Weakness	152	87.9%
Waegeneers, 1993 398	100 Hands	Weakness	43	43.0%
Pagnanelli, 1992 400	456 Hands	Weakness	210	46.1%
Richman, 1989	12	Weakness	7	58.3%
Kulick, 1986 418	167	Weakness	20	12.0%
Freshwater,				
1978 426	22	Weakness	17	77.3%

Figure 19. Symptoms of patients with carpal tunnel syndrome

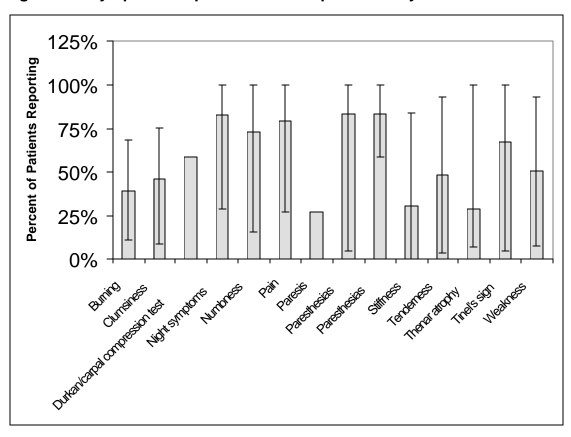


Table 49. Duration of symptoms among patients treated with surgery for carpal tunnel syndrome

Trial	N	Is duration of condition reported as Mean or Median?	Duration of condition before treatment (months)	Shortest period of duration before treatment (months)	Longest period of duration before treatment (months)
Porras, 2000 313	85	Mean	39	6	300
Straub, 1999 305	67	Median	24	3	300
Buchhorn, 1998 ³²⁸	50	Mean	43	Not reported	Not reported
Lee, 1998 331	525	Mean	40.1	2	480
Atroshi, 1997 337	204	Mean	24	1	240
Karlsson, 1997 ⁴⁸	74	Median	6	1	60
Leinberry, 1997 ³⁴²	44	Mean	31.8	3	168
Wheatly, 1997	126	Mean	90	10	120
Gibbs, 1996 351	46	Mean	57.0	1	360
Glowacki, 1996 ³⁵²	96	Mean	17.8	Not reported	Not reported
Lee, 1996 430	525	Mean	40.1	2	480
Nagle, 1996 357	506	Mean	31	1	420
Wintman, 1996 ³⁶²	50	Mean	28	3	173
Worseg, 1996	126	Mean	23.4	Not reported	Not reported
Mirza, 1995 371	236	Mean	23	Not reported	Not reported
Nancollas, 1995 ³⁷²	93	Mean	26.5	1	300
Sennwald, 1995 ³⁷³	47	Mean	9.2	Not reported	Not reported
Erdmann, 1994 ³⁰⁴	96	Mean	24.1	Not reported	Not reported
Roth, 1994 383		Mean	46.8	4	300
Brown, 1993 45	145	Mean	25	2	120
Clarke, 1993 431	37	Mean	37	2	300
Levine, 1993 393	39	Median	18	3	58
Palmer, 1993 397	173	Mean	35.6	Not reported	Not reported
Pagnanelli, 1992 400	228	Mean	45.6	3	360
Yu, 1992 403	53	Median	6	0	72

Trial	N	Is duration of condition reported as Mean or Median?	Duration of condition before treatment (months)	Shortest period of duration before treatment (months)	Longest period of duration before treatment (months)
Flaschka, 1991 ⁴⁰⁴	99	Mean	24	1	180
Hagberg, 1991 308	41	Mean	43.6	Not reported	Not reported
Jakab, 1991 406	73	Mean	48	2	516
Resnick, 1991	65	Mean	16.8	1	204
Richman, 1989 412	12	Mean	28	5	72
Szabo, 1989 284	22	Mean	29	7	120
Kulick, 1986	167	Mean	30	0	348
Shurr, 1986 420	36	Mean	12	Not reported	Not reported
Freshwater, 1978 ⁴²⁶	11	Mean	12	3	120

Table 50. Electrodiagnostic criteria among patients treated with surgery for carpal tunnel syndrome

Trial	Electrodiagnostic criteria
Hasegawa, 1999 ³²⁰	Patients with grade I (mild) symptoms were accepted for surgery if they also had distal motor latency >7.1ms or distal motor latency >5.2ms and 3 months of failed conservative treatment
Hirooka, 1999 ³²¹	Patients with grade 1 (mild) symptoms received surgery only if they had a distal motor latency of at least 7.0 ms.
Aulisa, 1998 ³²⁷	Patients fit into one of the following categories:
	Mild: Sensory conduction velocity, first digit to wrist <42m/s, third digit to wrist <44m/s Moderate: Sensory conduction velocity as in mild, plus median distal motor latency >4ms Severe: Absent sensory or motor median response.
Jacobsen, 1996 ³⁵³	Patients fit into one of the following categories:
	Slight CTS: >3 sensory responses delayed 2-4 standard deviations (SD). Intermediate CTS: All sensory responses delayed >3SD+decreased sensory amplitudes. Pronounced CTS: Several or all sensory responses lacking and rest are delayed >4SD with low amplitudes, motor delay >4SD with low amplitude or no motor response.
	The "normal" values to which these diagnostics were compared, and the size of a standard deviation were not reported.
Cook, 1995 432	Distal motor latency >4.5 ms and/or sensory antidromic latency >3.5 ms.
Lang, 1995 ¹⁰⁹	Either distal motor latency >4.5 ms or orthodromic sensory conduction velocity palm-to-wrist <45 m/s
Foulkes, 1994 376	Distal sensory latency of at least 3.6ms or motor latency of 4.4ms were considered supportive of diagnosis.
Pascoe, 1994 381	Difference between median and palmar sensory latency of more than 0.4ms
Brown, 1993 ⁴⁵	Electrophysiological confirmation was established when distal motor latency was 4.5 ms or there was a difference of 1 ms or more between the affected and unaffected hand or sensory latency was more than 3.5 ms or there was a difference of more than 0.5 ms between the affected and unaffected hand.
Nakamichi, 1993 ³⁹⁴	Distal motor latency >4.2ms or sensory nerve conduction velocity <45ms
Hagberg, 1991 308	A positive phalen test or distal motor latency of at least 4.5
Schuind, 1990 409	Distal motor latency >4ms or distal sensory latency >3.5ms
Richman, 1989 412	Distal motor latency >4.5ms or distal sensory latency >3.5ms
Szabo, 1989 ²⁸⁴	Distal motor latency >4.5 ms or distal sensory latency >3.5 ms.
Lowry, 1998 ⁴²⁹	Distal antidromic sensory latency >5ms or unobtainable at 13cm.
Holmgren-Larssen, 1985 433	Sensory nerve conduction velocity <50 ms and distal latency >4.5 ms.
Rhoades, 1985 422	Fibrillations in the abductor pollicis or opponens pollicis muscles detectable by EMG.
Van Rossum, 1980 424	Distal motor latency >4.5 ms
Freshwater, 1978 ⁴²⁶	No patients had normal motor latency (4.5ms or less), but this was not stated to have been an inclusion criterion.

Table 51. Reported occupations of patients receiving surgery for carpal tunnel syndrome

Study	Occupation	Number of Patients	Number of patients with occupation	Percent of patients with occupation
Mirza, 1995 371	Blue Collar	56	9	16.1%
Olney, 1999 323	Clerical	211	89	42.2%
Weber, 1997 347	Clerical	74	29	39.2%
Cobb, 1996 348	Clerical	235	38	16.2%
Mirza, 1995 371	Clerical	56	6	10.7%
Kelly, 1994 378	Clerical	69	10	14.5%
Palmer, 1993 397	Clerical	173	35	20.2%
Pagnanelli, 1992 400	Clerical	228	71	31.1%
Dumontier, 1995 365	Clerical, unoccupied or retired	96	47	49.0%
Wintman, 1996 362	Disabled	50	1	2.0%
Worseg, 1996 44	Employee	126	19	15.1%
Buchhorn, 1998 328	Employee- average work	50	21	42.0%
Olney, 1999 323	Factory	211	30	14.2%
Nagle, 1996 357	Heavy work	506	27	5.3%
Yu, 1992 ⁴⁰³	Heavy work	53	23	43.4%
Porras, 2000 313	High manual activity	85	14	16.5%
Kelly, 1994 378	High manual activity	69	7	10.1%
Cobb, 1996 348	Homemaker	235	19	8.1%
Wintman, 1996 362	Homemaker	50	12	24.0%
Worseg, 1996 44	Homemaker	126	8	6.3%
Mirza, 1995 371	Homemaker	56	5	8.9%
Chow, 1994 42	Homemaker	815	63	7.7%
Kelly, 1994 378	Homemaker	69	14	20.3%
Yu, 1992 ⁴⁰³	Homemaker	53	3	5.7%
Palmer, 1993 397	Industrial	173	90	52.0%
Katz, 1997 302	Laborer/machine operator	135	25	18.5%
Nagle, 1996 357	Light work	506	72	14.2%
Buchhorn, 1998 328	Light work	50	16	32.0%
Yu, 1992 ⁴⁰³	Light work	53	8	15.1%
Wintman, 1996 ³⁶²	Light labor with repetitive tasks or clerical work	50	15	30.0%
Nagle, 1996 357	Light-repetitive work	506	42	8.3%
Porras, 2000 313	Low manual activity	85	37	43.5%
Kelly, 1994 378	Low manual activity	69	21	30.4%
Katz, 1997 302	Management	135	22	16.3%
Weber, 1997 347	Management	74	14	18.9%
Lindau, 1999 322	Manual Worker	140	29	20.7%
Buchhorn, 1998 328	Manual Worker	50	8	16.0%
Weber, 1997 347	Manual Worker	74	25	33.8%
Cobb, 1996 348	Manual Worker	235	60	25.5%
Dumontier, 1995 365	Manual Worker	96	45	46.9%
Erhard, 1999 318	Manual worker- heavy lifting	124	12	9.7%
Olney, 1999 323	Manual worker- heavy lifting	211	40	19.0%

Study	Occupation	Number of Patients	Number of patients with occupation	Percent of patients with occupation
Buchhorn, 1998 328	Manual worker- heavy lifting	50	5	10.0%
Wintman, 1996 362	Manual worker- heavy lifting	50	5	10.0%
Chow, 1994 42	Manual worker- heavy lifting	815	322	39.5%
Pagnanelli, 1992 400	Manual worker- heavy lifting	228	60	26.3%
Erhard, 1999 318	Manual worker- light lifting	124	12	9.7%
Chow, 1994 42	Manual worker- light lifting	815	215	26.4%
Pagnanelli, 1992 400	Manual worker- light lifting	228	97	42.5%
Olney, 1999 323	Meat packing	211	15	7.1%
Palmer, 1993 397	Medical	173	7	4.0%
Porras, 2000 313	Medium manual activity	85	35	41.2%
Nagle, 1996 357	Medium work	506	46	9.1%
Yu, 1992 ⁴⁰³	Medium strenuous work	53	13	24.5%
Lindau, 1999 322	Nonmanual worker	140	41	29.3%
Chow, 1994 42	Other	815	68	8.3%
Katz, 1997 302	Other	135	81	60.0%
Cobb, 1996 348	Other	235	14	6.0%
Worseg, 1996 44	Other	126	3	2.4%
Kelly, 1994 378	Other	69	1	1.4%
Palmer, 1993 397	Other	173	15	8.7%
Wintman, 1996 362	Professional	50	6	12.0%
Mirza, 1995 371	Professional	56	11	19.6%
Palmer, 1993 397	Professional	173	16	9.2%
Palmer, 1993 397	Education	173	8	4.6%
Lindau, 1999 322	Retired	140	21	15.0%
Weber, 1997 347	Retired	74	6	8.1%
Wintman, 1996 362	Retired	50	7	14.0%
Worseg, 1996 44	Retired	126	60	47.6%
Hallock, 1995 368	Retired	100	15	15.0%
Mirza, 1995 371	Retired	56	5	8.9%
Strasberg, 1994 386	Retired	45	4	8.9%
Yu, 1992 ⁴⁰³	Retired	53	6	11.3%
Palmer, 1993 397	Retired or Homemaker	173	40	23.1%
Olney, 1999 323	Retired or light employment	211	57	27.0%
Chow, 1994 42	Retired or unemployed	815	147	18.0%
Kelly, 1994 378	Retired or unemployed	69	16	23.2%
Erhard, 1999 318	Sedentary	124	18	14.5%
Nagle, 1996 357	Sedentary	506	69	13.6%
Strasberg, 1994 ³⁸⁶	Student	45	2	4.4%
Wintman, 1996 362	Unemployed	50	4	8.0%
Worseg, 1996 44	Unemployed	126	19	15.1%
Strasberg, 1994 386	Unemployed	45	28	62.2%
Worseg, 1996 44	Worker	126	17	13.5%

Table 52. Excluded trials

Study	Reason for Exclusion
Todnem, 2000 ⁴³⁵	Retrospective comparison of operated and nonoperated patients. Groups were significantly different in several electrodiagnostic parameters prior to surgery.
Atherton, 1999 436	Did not report any patient characteristics or patient-oriented outcomes.
Brüser, 1999 437	A single study comparing two very similar treatments.
Davis, 1998 438	Utilized a combination of treatments, rendering it impossible to determine the effect of a single treatment.
Ebenbichler, 1998 ⁴³⁹	There were significant differences between groups at baseline. Although patients were described as having bilateral carpal tunnel syndrome, five patients in the treated group and seven in the placebo group had no wrist complaints.
Garfinkel, 1998 440	The treatment received by the control group was not standardized and was not described.
Netscher, 1998 47	Did not report any patient-oriented outcomes.
Rozmaryn, 1998 32	Patients received an assortment of nonstandardized treatments in addition to the experimental treatment.
Braithwaite, 1997 441	Compares minor variations in surgical technique. No patient-oriented outcome measures were reported other than perioperative pain. No patient characteristics were reported.
Jones, 1997 442	A single study comparing two very similar treatments.
Monge, 1995 ⁴⁴³	No patient-oriented outcomes were reported for the controls; only for treated patients. Reported no information on the source of control data or the comparability of controls and treated patients.
Bande, 1994 444	Groups were not comparable. Patients with comorbidities (e.g. synovitis, diabetes, rheumatoid arthritis) were all placed in the open release group. There was no indication as to how many such patients were included.
Biyani, 1993 388	A single study comparing two very similar treatments.
Nathan, 1993 395	A single study comparing two very similar treatments.
Spooner, 1993 445	Did not report any patient-oriented outcomes.
Groves, 1989 446	Compared outcomes at two separate clinics. There was no indication that the patient populations treated by the two clinics were comparable. This study had no internal validity.
Wolaniuk, 1983 447	Did not report any patient-oriented outcomes.
Ellis, 1979 447	Describes a double-blind crossover study of a single patient.

Table 53. Internal validity of studies comparing open and endoscopic carpal tunnel release

Study	Number of patients	Percent of patients with bilateral procedures	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?
Concannon, 2000 ⁴⁴⁹	191	NR ^a	Single	Not reported	Retro	No	0	Yes
Chen, 1999 317	948	At least 34.8% ^b	Single	Not reported	Retro	No	24	No
Hasegawa, 1999 ³²⁰	82	2.4%	Single	Not reported	Retro	No	0	Yes
Povlsen, 1997 ⁴⁵⁰	120	0%	Multiple (<5)	Not reported	СТ	No	4	No
Gibbs, 1996 351	46	23.9%	Single	Not reported	Retro	No	3	No
Jacobsen, 1996 ³⁵³	29	10.3%	Single	Not reported	RCT	Rater	0	Yes
Worseg, 1996 ⁴⁴	126	0%	Single	Not reported	СТ	No	0	Yes
Dumontier, 1995 ³⁶⁵	103	0%	Single	Not reported	RCT	No	83	No
Futami, 1995 366	10	100%	Single	Not reported	СТ	No	0	Yes
Hallock, 1995 ³⁶⁸	96	37%	Single	Not reported	СТ	No	0	Noc
Sennwald, 1995 ³⁷³	47	0%	Single	Not reported	RCT	No	0	Yes
Erdmann, 1994 ³⁰⁴	71	47.9%	Single	Not reported	RCT	No	0	Yes
Kerr, 1994 379	157	At least 17.4% ^b	Single	Not reported	СТ	No	13	No
Brown, 1993	151	13.2%	Multiple (<5)	No	RCT	Rater	22	No
McDonough, 1993 448	88	23.5%	Single	Yes	Retro	No	7	No
Palmer, 1993 ³⁹⁷	211	29.4%	Single	No	СТ	No	0	Yes
Agee, 1992	122	20.5%	Multiple (>5)	Yes	RCT	No	NR	No

a: This report describes the results of 191 procedures. The number of patients was not reported.
b: The number of bilateral procedures among those patients who underwent open procedures was not reported.
c: Four patients whose endoscopic procedures were, for various technical reasons, converted to open procedures, are included in the Open group.

Table 54. Generalizability of studies comparing open and endoscopic carpal tunnel release

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Concannon, 2000 ⁴⁴⁹	191	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	No	No
Chen, 1999 317	948	48 (21-79)	78.5	NR	0.6	2.4	0	NR	0	0.3	0	Yes	No
Hasegawa, 1999 ³²⁰	82	54.1	100	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes
Povlsen, 1997 450	120	NR	NR	NR	NR	NR	NR	NR	0	NR	NR	No	No
Gibbs, 1996 351	46	56.2 (31-86)	89.1	57.0 (1-360)	0	0	NR	0	NR	NR	NR	No	No
Jacobsen, 1996 353	29	(24-59)	NR	NR	NR	NR	NR	NR	NR	NR	NR	Yes	Yes
Worseg, 1996 44	126	56.0 (35-90)	69.8	23.4	NR	0	NR	NR	0	NR	NR	Yes	Yes
Dumontier, 1995 365	103	52.3	82.5	NR	NR	NR	NR	NR	NR	NR	NR	No	No
Futami, 1995 366	10	53 (39-61)	90.0	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes
Hallock, 1995 368	96	44.2	77.1	NR	NR	NR	0	NR	NR	NR	NR	No	No
Sennwald, 1995 373	47	52.5	80.9	9.2	0	0	NR	NR	NR	NR	NR	No	Yes
Erdmann, 1994 304	71	53.4	98.6	27.3	2.8	28.2	0	NR	NR	NR	NR	Yes	No
Kerr, 1994 379	157	44.8 (19-82)	56.5	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
	151	55 (25-87)	65.6	25 (2-120)	NR	0	NR	NR	0	0	NR	No	No
McDonough, 1993 ⁴⁴⁸	88	46.0 (21-79)	62.5	35.6	NR	NR	NR	NR	NR	NR	NR	No	Yes
Palmer, 1993 397	211	44.9 (20-83)	65.4	35.7	1.4	0	NR	NR	NR	NR	NR	No	Yes

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Agee, 1992 46	122	NR	NR	NR	0	0	0	NR	0	NR	0	Yes	Yes

Table 55. Patient employment characteristics in studies comparing open and endoscopic carpal tunnel release

Study	Number of patients	% Patients employed	% Patients receiving workers' compensation	% Patients retired	% Patients Homemakers	Reported occupations
Concannon, 2000 ⁴⁴⁹	191	Not reported	44.0	Not reported	Not reported	Not reported
Chen, 1999 317	948	Not reported	Not reported	Not reported	Not reported	Not reported
Hasegawa, 1999 ³²⁰	82	Not reported	Not reported	Not reported	Not reported	Not reported
Povlsen, 1997 450	120	Not reported	Not reported	Not reported	Not reported	Not reported
Gibbs, 1996 351	46	84.8	15.2	Not reported	Not reported	16 Retired, homemaker or unemployed
Jacobsen, 1996 ³⁵³	29	100	0	0	0	Not reported
Worseg, 1996 44	126	31.0	87.3	47.6	6.3	19 Employee 17 Worker 60 Retired 19 Unemployed 8 Homemaker 3 Other
Dumontier, 1995 ³⁶⁵	103	89.3	Not reported	Not reported	Not reported	45 Manual workers 47 Clerical, unoccupied or retired
Futami, 1995	10	Not reported	Not reported	Not reported	Not reported	Not reported
Hallock, 1995 368	96	Not reported	54.2	15.6	Not reported	Not reported
Sennwald, 1995 ³⁷³	47	Not reported	Not reported	Not reported	Not reported	Not reported
Erdmann, 1994 ³⁰⁴	71	Not reported	Not reported	Not reported	Not reported	Not reported
Kerr, 1994 ³⁷⁹	157	Not reported	Not reported	Not reported	Not reported	Not reported
Brown, 1993 45	151	53.6	4.6	Not reported	Not reported	41 Professional, management or business 29 Clerical or technical support 11 Manual labor
McDonough, 1993 ⁴⁴⁸	88	Not reported	27.3	Not reported	Not reported	Not reported

Study	Number of patients	% Patients employed	% Patients receiving workers' compensation	% Patients retired	% Patients Homemakers	Reported occupations
Palmer, 1993 ³⁹⁷	211	73.9	57.8	Not reported		8 Education 90 Industrial 7 Medical 16 Professional 35 Clerical 40 Retired or Homemaker 15 Other
Agee, 1992 46	122	Not reported		Not reported	Not reported	Not reported

Table 56. Global outcome in patients treated with open or endoscopic carpal tunnel release

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Hasegawa et al., 1999 ³²⁰	40 Open 42 Endoscopic	Global outcome rating at 12 Months 28 Excellent 8 Good 3 Fair 1 Poor 29 Excellent 13 Good 1 Fair 1 Poor	Not significantly different by chi square test conducted by ECRI, p = 0.57
Gibbs et al., 1996 ³⁵¹	43 Open 14 Endoscopic (Hands)	Mean change in symptom severity score 3-33 Months -12.5±5.6 -12.2±5.3	Not significantly different by t test conducted by ECRI, p = 0.86
Worseg et al., 1996 ⁴⁴	62 Open 64 Endoscopic	Mean symptom rating, verbal scale This outcome was reported using a 3-dimensional graph, making it difficult to estimate values.	Scores were not significantly different between groups at any time point (p >0.05, Wilcoxon rank sum test)
Futami 1995 366	10 Open 10 Endoscopic (Hands of 10 patients)	Weeks until relief of symptoms 2.5 Weeks 2.4 Weeks	Not reported
Hallock 1995 ³⁶⁸	71 Open 66 Endoscopic (Hands)	Number of hands with complete relief of symptoms (Time not reported) 63 61	Not significantly different by chi square test conducted by ECRI, p = 0.46
Erdmann, 1994 ³⁰⁴	52 Open 53 Endoscopic	Days until relief of symptoms 1.75 Days 1.1 Days	Not significantly different by Mann-Whitney test. The p value determining significance was not reported.
Brown, 1993 ⁴⁵	82 Open 78 Endoscopic (Hands)	Mean patient satisfaction rating, 0-100 84 Days: 84±26 84 Days: 89±18	Not significantly different by t test conducted by ECRI, p = 0.15

Table 57. Results of meta-analysis of the effect of open or endoscopic treatment on global outcome

Author	Year	N	Effect Size	95% CI	p-value	Standardize d Residual	Outlier by Std Resid?
Hasegawa ³²⁰	1999	82	0.362	-0.07-0.80	0.105	0.83	No
Gibbs 351	1996	57	-0.054	-0.66-0.55	0.862	-0.84	No
Worseg 44	1996	126	0.12a	-0.23-0.41.	0.502	-0.49	No
Hallock ³⁶⁸	1995	137	0.240	-0.41-0.89	0.466	0.15	No
Brown 45	1993	160	0.222	-0.09-0.53	0.163	-0.22	No
			Effect Size	05% CI	n value	O Statistic	n of O

	Summary Effect Size	95% CI	p-value	Q Statistic	p of Q
Fixed Effects Model	0.19	0.01-0.38	0.041	1.44	0.838

 $^{^{\}mathrm{a}}$: Estimated from published data by assuming that the pvalue of the Wilcoxon test was 0.5

Table 58. Time to return to work in patients treated with open or endoscopic surgery

Study	n (units)	Time Until Return to Work	Statistical Significance of Difference Between Groups
Gibbs, 1996 351		Time at which 50% of patients had returned to work	Groups were not significantly different by log rank test, p = 0.63
	Open	4 Days (Range 1->1003) ^a	
	Endoscopic	14 Days Range (1-91)	
	Total N = 28 Group n not reported		
Jacobsen, 1996 ³⁵³	16 Open	Open 18.94±10.25 Days (Range 0-42)	Groups not significantly different, p >0.05, Fisher Exact test
	16 Endoscopic (Hands)	Endoscopic 17.06±9.11 Days (Range 0-31)	
Dumontier, 1995 ³⁶⁵		Percent of patients returning to work within:	Groups were not significantly different at any time by chi square test.
	Open	2 Weeks: 29%; 1 Month: 70%; 3 Months: 89% ^b	At 1 month, p = 0.13. p-values were not reported for the other two time points.
	Endoscopic	2 Weeks: 30%; 1 Month 45%; 3 Months 70%	
	Numbers of patients not reported	o monus 7070	
Futami, 1995 ³⁶⁶	Open 3	7 Weeks	Not reported
	Endoscopic 3	6 Weeks	
Hallock, 1995 ³⁶⁸	Open 39	46.3±36.9 Days ^c	Groups were not significantly different, p = 0.373. The test used was not
	Endoscopic 25	39.8±19.3 Days	reported.
Sennwald, 1995 ³⁷³	22 Open	41.95±13.18 Days ^d	Groups were significantly different by test calculated by ECRI, p = 0.000001
	25 Endoscopic	24.13±7.69 Days	
	(Patients)		
Erdmann, 1994 ³⁰⁴	23 Open (Patients) 27 Open (Hands) ^e	39 Days Open	Groups were significantly different, p <0.005 unpaired Mann-Whitney U test
	23 Endoscopic (Patients) 28 Endoscopic (Hands)	14 Days Endoscopic	

Study	n (units)	Time Until Return to Work	Statistical Significance of Difference Between Groups
Kerr, 1994 379	72 Open	Patients treated endoscopically returned to work 10.6 days	Groups were significantly different by paired t-test (p = 0.0015)
	72 Endoscopic	sooner than those treated openly.	
Brown, 1993 ⁴⁵	85 Open	Median 28 Days Open ^a	Groups were statistically significant, p <0.05, log-rank test
	84 Endoscopic (Hands)	Median 14 Days Endoscopic	
McDonough, 1993 448	28 Open	50.4 Days (Range 11-103)	Not reported
	27 Endoscopic (Patients)	28.5 Days (Range 4-67)	
Palmer, 1993 ³⁹⁷	Open	44.1±37.3	Open was significantly different from the other two groups by t-test, p <0.05
	Endoscopic- Agee method	20.7±12.8	
	Endoscopic- Chow method	27.9±16.9	
	n not reported		
Agee, 1992 ⁴⁶	30 Open	Median 46.5 Days ^a	Statistically significant difference between groups, p <0.01, survival
	49 Endoscopic (Patients)	Median 25 Days	analysis version of the Wilcoxon test

a: Calculated by Kaplan-Meier survival analysis b: Percentages estimated from a published chart. They cannot be converted to numbers of patients because it is unclear whether they are percentages of all patients or of patients employed prior to surgery.

c: Some patients in each group did not return to work. The numbers reported therefore do not constitute an accurate representation

of time to return to work.

d: Estimated by ECRI from a published chart.

e: Unclear whether data is reported per patient or per treated hand. Therefore, we did not calculate an effect size for this study.

Table 59. Summary of effect of treatment type on return to work

Dumontier, 1995 365	Study	Which Procedure Yielded Faster Return to Work?	Was the Difference Statistically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Dumontier, 1995 3/65	Gibbs, 1996 351	Open	No	Not calculable	Not calculable
2 weeks Open at 1 month and 3 months Futami, 1995 366 Endoscopic No Not calculable Not calculable Not calculable Sennwald, 1995 373 Endoscopic Yes 15.1% 1.65 (0.99 –2.31) Erdmann, 1994 304 Endoscopic Yes Not calculable Not calculable	Jacobsen, 1996 ³⁵³	Endoscopic	No	25%	0.19 (-0.51 – 0.88)
Futami, 1995 366 Endoscopic No Not calculable Not calculable Hallock, 1995 368 Endoscopic No 32.6% Not calculable Sennwald, 1995 373 Endoscopic Yes 15.1% 1.65 (0.99 –2.31) Erdmann, 1994 304 Endoscopic Yes Not calculable Not calculable Kerr, 1994 379 Endoscopic Yes Not calculable Not calculable Brown, 1993 45 Endoscopic Yes Not calculable Not calculable McDonough, 1993 Endoscopic Not reported Not calculable Not calculable	Dumontier, 1995 ³⁶⁵	2 weeks Open at 1 month	No	Not calculable	Not calculable
Hallock, 1995 368EndoscopicNo32.6%Not calculableSennwald, 1995 373EndoscopicYes15.1%1.65 (0.99 -2.31)Erdmann, 1994 304EndoscopicYesNot calculableNot calculableKerr, 1994 379EndoscopicYesNot calculableNot calculableBrown, 1993 45EndoscopicYesNot calculableNot calculableMcDonough, 1993EndoscopicNot reportedNot calculableNot calculable	Futami 1995 366		No	Not calculable	Not calculable
Sennwald, 1995 373EndoscopicYes15.1%1.65 (0.99 –2.31)Erdmann, 1994 304EndoscopicYesNot calculableNot calculableKerr, 1994 379EndoscopicYesNot calculableNot calculableBrown, 1993 45EndoscopicYesNot calculableNot calculableMcDonough, 1993EndoscopicNot reportedNot calculableNot calculable					
Erdmann, 1994 304 Endoscopic Yes Not calculable Not calculable Kerr, 1994 379 Endoscopic Yes Not calculable Not calculable Brown, 1993 45 Endoscopic Yes Not calculable Not calculable McDonough, 1993 Endoscopic Not reported Not calculable Not calculable					
Kerr, 1994 379EndoscopicYesNot calculableNot calculableBrown, 1993 45EndoscopicYesNot calculableNot calculableMcDonough, 1993EndoscopicNot reportedNot calculableNot calculable		•			
Brown, 1993 ⁴⁵ Endoscopic Yes Not calculable Not calculable McDonough, 1993 Endoscopic Not reported Not calculable Not calculable					
McDonough, 1993 Endoscopic Not reported Not calculable Not calculable					
Palmer 1993 397 Endoscopic Yes Not calculable Not calculable	McDonough, 1993				
Trainior, 1770 Endoscopio 165 International International	Palmer, 1993 397	Endoscopic	Yes	Not calculable	Not calculable
Agee, 1992 46 Endoscopic Yes Not calculable Not calculable		•			

a: Calculated by ECRI

Table 60. Time to return to activities of daily living in patients treated with open or endoscopic surgery

Study	Number of Patients	Time to Return to Activities of Daily Living	Statistical Significance of Difference Between Groups
Gibbs, 1996 351		Time until 50% of patients had returned to ADL ^a	Groups not significantly different by log-rank test
	43 Open	21 Days (Range 1->911)	
	14 Endoscopic	21 Days (Range 7->425)	
Futami, 1995 ³⁶⁶	10 Open ^b	41 Days (Range 28-51)	Groups significantly different by t-test, p < 0.01
	10 Endoscopic	12 Days (Range 4-18)	
Erdmann, 1994 ³⁰⁴	23 Open (Patients) 27 Open (Hands)	39 Days	Groups significantly different (p <0.005, Mann-Whitney test)
	23 Endoscopic (Patients) ^c 28 Endoscopic (Hands)	14 Days	
Brown, 1993 ⁴⁵	21 Days, N = 149 Hands Group n not reported ^d	Number of patients (hands) with no impairment of ADL	
	Open	3 (5)	Groups were not significantly different by Kaplan-Meier survivorship analysis.
	Endoscopic	8 (8)	Sa meremp analysis
	42 Days, N = 147 Hands		
	Open	(12)	Groups were not significantly different by Kaplan-Meier
	Endoscopic	(14)	survivorship analysis.
	84 Days, N = 160 Hands		Crounc ware not cignificantly
	82 Open	28 (29)	Groups were not significantly different by Kaplan-Meier survivorship analysis. However,
	78 Endoscopic	39 (42)	they were significantly different by chi square test conducted by ECRI, p = 0.019

Study	Number of Patients	Time to Return to Activities of Daily Living	Statistical Significance of Difference Between Groups
Agee, 1992 46	63 Open	Median 13 Days, estimated by Kaplan-Meier	Groups not significantly different according to a survival analysis
	81 Endoscopic		version of the Wilcoxon test.
		Median 9 Days, estimated	
	(Hands)	by Kaplan-Meier	

Table 61. Summary of effect of treatment (open or endoscopic) on time to return to ADLs

Study	Which Procedure Yielded Faster Return to Daily Activities?	Was the Difference Statistically Significant?	Power (Minimum percent difference detectable	Effect Size (95% Confidence Interval) ^a
Gibbs, 1996 351	Both groups were equal	No	Not calculable	Not calculable
Futami, 1995 366	Endoscopic	Yes	Not calculable	Not calculable
Erdmann, 1994 304	Endoscopic	Yes	Not calculable	Not calculable
Brown, 1993 ⁴⁵	Endoscopic	21 days: No 42 days: No 84 days: Yes	21 days: Not calculable 42 days: Not calculable 84 days: 18.3%	21 Days: Not calculable 42 days: Not calculable 84 days: 0.42 (0.065-0.77)
Agee, 1992 46	Endoscopic	No	Not calculable	Not calculable

a: Calculated by ECRI

a: Calculated by Kaplan-Meier survival analysis
 b: 20 hands in 10 patients
 c: Unclear whether means were calculated as per patient or per hand.
 d: Sum of group ns calculated by ECRI from published data did not match reported total Ns.

Table 62. Symptomatic pain in patients treated with open or endoscopic carpal tunnel release

Study	Number of Hands	Pain	Statistical Significance of Difference Between Groups
Gibbs, 1996 351		Pain rating	Groups not significantly different
	43 Open	Preop:: 3.3±1.0 18.9 Months: 1.2 ±0.52	before or after treatment by t test, p = 0.78 and 0.21 respectively.
	14 Endoscopic	Preop: 3.3±0.87 16 Months: 1.5±0.96	
Erdmann, 1994 ³⁰⁴		Mean VAS, 0-10 Scale ^a	Groups significantly different at 1 week only (Mann-Whitney
	52 Open	Preop: 5.6; 1 Week: 3.9 1 Year: 0.95	test, p <0.05)
	53 Endoscopic	Preop, 5.7; 1 Week: 2.4 1 Year: 0.1	
Palmer, 1993 ³⁹⁷		Percent of patients ^b reporting nocturnal pain	Groups not significantly different at any time point by chi square test, p >0.05
	42 Patients, 49 Hands Open	Preop: 88.7% 2 Weeks: 23.3% 6 Months: 25.0%	
	70 Patients, 90 Hands Endoscopic (Agee method)	Preop: 80.0% 2 Weeks: 16.7% 6 Months: 12.5%	
	62 Patients, 72 Hands Endoscopic (Chow method)	Preop: 89.8% 2 Weeks: 21.7% 6 Months: 28.9%	
Agee, 1992 46		Percent of patients ^b with symptomatic pain	Not reported
	65 Open	Preop: 89; 1 Week: 59 26 Weeks: 27	
	82 Endoscopic	Preop: 85; 1 Week: 43 26 Weeks: 25	

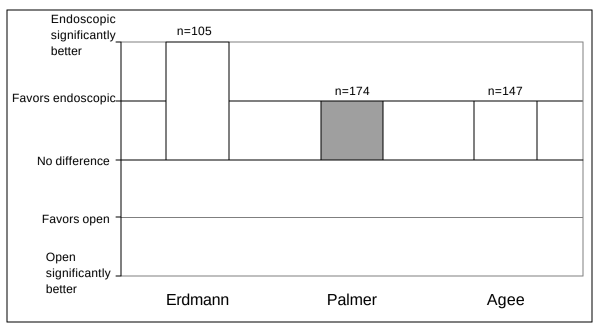
a: Estimated by ECRI from a published chart
b: The report states that outcomes are reported as percent of patients. However, as some patients had a different procedure in each hand, it is likely that the outcome is actually percent of hands. Thus, the true n is unclear.

Table 63. Summary of effect of treatment (open or endoscopic) on pain

Study	Which Procedure Had Less Pain?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable 80% of the time) ^a	Effect Size (95% Confidence Interval) ^a
Gibbs, 1996 ³⁵¹	Preop: No difference Early: Not reported Late: Open	Preop: No Early: Not reported Late: No	Preop: 17.5% Early: Not reported Late: 40.0%	Preop: 0.0 (-0.60–0.60) Early: Not reported Late: -0.45 (-1.06-0.15)
Erdmann, 1994 ³⁰⁴	Preop: Open Early: Endoscopic Late: Endpscopic	Preop: No Early: Yes Late: No	Not calculable	Preop: Not calculable Early: 0.39 (0.00-0.77) ^b Late: Not calculable
Palmer, 1993 ³⁹⁷	Preop: Endoscopic Early: Endoscopic Late: Endoscopic	Preop: No Early: No Late: No	Not calculable	Not calculable
Agee, 1992	Preop: Endoscopic Early: Endoscopic Late: Endoscopic	Preop: Not reported Early: Not reported Late: Not reported	Not calculable	Not calculable

a: Calculated by ECRI

Figure 29. Summary of the effect of treatment on pain at early time points



An open bar indicates an RCT, a striped bar a CT. The study by Gibbs does not appear because it did not report early time ponts.

b: Calculated by ECRI based on the conservative assumption that p = 0.049)

Function

Function refers to the ability of the patient to perform various tasks and activities with their affected limb(s). It is measured using any of a number of tests.

Only one nonrandomized controlled trial, that by Worseg, reported a measure of function. This outcome is described in Table 64 and summarized in Table 65. Worseg's global function was the mean of the difficulty ratings (scale of 1-5) of eight individual activities (writing, buttoning clothes, holding a book, gripping a telephone, opening jars, household chores, carrying a grocery bag, and bathing and dressing).

The endoscopic group experienced superior function one week after surgery, but there were no statistically significant differences in the long term. This is consistent with the idea that the less invasive treatment leads to more rapid recovery. Because, however, function was examined in only one study (which was not randomized), it is difficult to draw firm evidence-based conclusions about the relative effects of open and endoscopic surgery on function.

Table 64. Function in patients treated with open or endoscopic carpal tunnel release

Study	Number of Patients	Function	Statistical Significance of Difference Between Groups	
Worseg et al., 1996 ⁴⁴	Open 62	Mean of function scores ^a Preop: 3.14; 1 Week: 3.33; 24 Weeks: 1.29	Between group differences were significant at 1 Week only (p <0.05, Wilcoxon rank sum test).	
	Endoscopic 64	Preop: 3.16; 1 Week: 2.29; 24 Weeks: 1.20		

a: Lower score indicates superior function

Table 65. Summary of the effect of treatment on function

Study	Which Procedure Had Superior Function at Followup?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Worseg, 1996 ⁴⁴	Endoscopic	At 1 week only	Not calculable	Preop: 0.12 (-0.23 – 0.47) ^b 1 Week: 0.35 (0.00 – 0.70) 24 Weeks: 0.12 (-0.23 – 0.47)

a: Calculated by ECRI

b: Calculated by ECRI based on the conservative assumption that p = 0.49 at one week and p = 0.50 at the other time points.

Table 66. Blood vessel, nerve and tendon lacerations during open and endoscopic carpal tunnel release

Study	Procedures	Endoscopic Lacerations	Open Lacerations
Chen, 1999 317	Open 64 Endo 1214	1 Motor nerve	0
Povlsen, 1997 450	Open 50 Endo 50	0	0
Jacobsen, 1996 353	Open 16 Endo 16	0	0
Worseg, 1996 44	Open 62 Endo 64	1 Transection of the superficial palmar arch	0
Dumontier, 1995 365	Open 40 Endo 56	1 Ulnar artery injury	0
Sennwald, 1995	Open 22 Endo 25	0	0
Erdmann, 1994 ³⁰⁴	Open 52 Endo 53	0	1 Palmar cutaneous nerve
Brown, 1993 45	Open 85 Endo 84	1 Superficial palmar arch	0
McDonough, 1993	Open 50 Endo 50	1 Digital tendon	0
Palmer, 1993 ³⁹⁷	Open 49 Endo (Agee) 90	0	0
	Endo (Chow) 72		
Total		1774 Procedures	490 Procedures
		5 Lacerations	1 Laceration

Table 67. Incomplete transections of the carpal ligament

Study	Procedures	Endoscopic Incomplete Transections	Open Incomplete Transections
Concannon et al., 2000 449	Open 103 Endo 88	5	0
Sennwald and Benedetti, 1995 373	Open 22 Endo 25	0	0
Erdmann, 1994 ³⁰⁴	Open 52 Endo 53	1	0
McDonough et al., 1993 448	Open 50 Endo 50	1	0
Palmer et al., 1993 397	Open 49 Endo (Agee) 90 Endo (Chow) 72	1 Agee 1 Chow	0
	Lituo (Chow) 72	I I Chow	
Т	otal	378 Procedures	276 Procedures
		9 Incomplete transections	0 Incomplete transections

Conclusions

Endoscopic release allows faster return to work and to activities of daily living. In addition, it leads to superior global outcome and reduced pain. However, the effects on pain and global outcome may be small. Presently available data do not allow one to reach firm evidence-based conclusions about the relative effect of open and endoscopic surgery on function. Because of incomplete transection of the transverse carpal ligament, endoscopic release has a higher rate of reoperation. Although there is insufficient data to draw firm conclusions, endoscopic release may also have a higher complication rate.

Table 68. Internal validity of studies comparing open carpal tunnel release with and without neurolysis

Study	Number of patients	Percent of patients with bilateral procedures	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?
Leinberry, 1997 ³⁴²	44	13.6%	Single	No	RCT	Rater	0	Yes
Blair, 1996 428	117	36.0%	Single	No	RCT	Rater	42	No
Foulkes, 1994 ³⁷⁶	46	8.7%	Single	No	RCT	Rater	23	No
Mackinnon, 1991 ⁴⁰⁷	59	6.8%	Single	No	RCT	Double	20	No
Lowry, 1988 429	50	22.0%	Single	Not reported	RCT	Double	3	No
Gelberman, 1987 415	61	13.1%	Multiple (<5)	No	Retro	No	0	Yes
Holmgren- Larsson, 1985 ⁴³³ Holmgren, 1987 ⁴¹⁶	48	0.0%	Single	Not reported	RCT	No	7	No
Freshwater, 1978 ⁴²⁶	22	18.2%	Single	Not reported	СТ	Double	0	Yes

Table 69. Generalizability of studies comparing open carpal tunnel release with and without neurolysis

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Leinberry, 1997	44	65 (38-100)	59.1	31.8 (1-360)	6.8	NR	NR	NR	NR	NR	NR	No	Yes
Blair, 1996 428	86	49 (23-82)	72.1	NR	NR	NR	NR	0	0	NR	NR	No	Yes
Foulkes, 1994 376	46	NR	37.0	NR	NR	NR	NR	NR	NR	0	NR	No	Yes
Mackinnon, 1991 ⁴⁰⁷	79	58.5 (20-91)	60.8	NR	0	NR	0	0	0	NR	NR	No	Yes
Lowry, 1988 429	50	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes
Gelberman, 1987 415	61	59.2 (28-90)	37.7	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes
Holmgren, 1987	48	50 (21-80)	68.8	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes
Holmgren- Larsson, 1985	48	50 (21-80)	68.8	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes
Freshwater, 1978 426	22	NR; (32-74)	NR	12 (3-120)	NR	0	NR	NR	NR	NR	NR	NR	NR

Table 70. Patient employment characteristics in studies comparing open carpal tunnel release with and without neurolysis

Study	Number of patients	% Patients employed	% Patients receiving workers' compensation	% Patients retired	% Patients Homemakers	Reported occupations
Leinberry, 1997 342	44	Not reported	Not reported	Not reported	Not reported	Not reported
Blair, 1996 428	86	Not reported	Not reported	Not reported	Not reported	Not reported
Foulkes, 1994 376	46	Not reported	Not reported	Not reported	Not reported	Not reported
Mackinnon, 1991	79	Not reported	12.7	Not reported	Not reported	Not reported
Lowry, 1988 429	50	Not reported	Not reported	Not reported	Not reported	Not reported
Gelberman, 1987	61	Not reported	Not reported	Not reported	Not reported	Not reported
Holmgren, 1987 416	48	Not reported	Not reported	Not reported	Not reported	Not reported
Holmgren-Larsson, 1985 ⁴³³	48	Not reported	Not reported	Not reported	Not reported	Not reported
Freshwater, 1978	22	Not reported	Not reported	Not reported	Not reported	Not reported

Table 71. Effect of neurolysis on global outcome

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Leinberry, 1997 342		Number of hands with no symptoms	Not significantly different, test not reported
	Open Release 25	12 Months: 15	
	Release and Neurolysis 25	12 Months: 14	
	(Hands)		
Blair, 1996 ⁴²⁸		Patients stating they would have surgery again	Not reported
	Open Release 27	26	
	Release and Neurolysis 48	46	
	(Hands)	Patient perceptions about relief of symptoms	
	Open Release 27	Permanent total: 13 Permanent partial: 12 Temporary total: 2	
	Release and Neurolysis 48	Permanent total: 31 Permanent partial:15 Temporary total: 2	
		Patient satisfaction	
	Open Release 27 Release and	Happy/very happy: 19 Satisfied, with reservations: 8 Disappointed/ very disappointed: 0	
	Neurolysis 48	Happy/very happy: 35 Satisfied, with reservations: 9 Disappointed/ very disappointed: 4	

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Foulkes, 1994 376		Improvement at 29 Months	Not reported
	Open Release 8	Normal 2 Improved 6 Unimproved 0	
	Release and Neurolysis 15	Normal 5 Improved 9 Unimproved 1	
	Recalculated: Open Release 10 ^a	Recalculated: Normal 2 Improved 6 Unimproved 2	Not reported
	Release and Neurolysis 26	Normal 5 Improved 9 Unimproved 12	
	(Hands)	Symptom severity score	
	Open Release 8	Preop: 2.5; 29 Months: 0.4 Recalculated to account for patient attrition:	
	Open Release 10	Preop: 2.5; 29 Months: 0.82	
	Release and Neurolysis 15	Preop 2.9; 29 Months: 0.3 Recalculated to account for patient attrition:	
	Release and Neurolysis 26	Preop: 2.9; 29 Months: 1.4	
	(Hands)		
Mackinnon 1991 407		Symptom rating at 12 months.	Not reported
	Open Release 32	Relief of all or most symptoms 28 Unimproved 4 Worse 0	
	Release and neurolysis 31	Relief of all or most symptoms 25 Unimproved 5 Worse 1	
	(Hands)		

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Lowry, 1988 429		3 Months	Not reported
	Open Release 23	Excellent 7 Good 8 Fair 6 Poor 2	
	Release and Neurolysis 24	Excellent 4 Good 12 Fair 7 Poor 1 Recalculatedb:	
	Open Release 25	Excellent 7 Good 8 Fair 6 Poor 4	
	Release and Neurolysis 25	Excellent 4 Good 12 Fair 7 Poor 2	
Gelberman,1987 ⁴¹⁵ ; Rhodes, 1985 ⁴⁵¹	Open Release: 29	Number of patients with complete resolution of signs and symptoms Complete resolution: 18 Mean followup time: 16 Months	Significantly different (p <0.05, chi square)
	Release and Neurolysis 32	Complete resolution: 10 Mean followup time: 18 Months	

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Holmgren-Larsson, et al. 1985 433	48 Patients; Number in each group not reported.	Percent of patients reporting themselves symptom-free at 6 months	Not reported
	Open Release	89%	
Holmgren, 1987 ⁴¹⁶	Release and Neurolysis	89%	
Hollingren, 1967 110		3-4 Years:	
	Open Release 20	Totally restituted: 12 Improved: 4 Dead: 1 Did not respond: 3	
	Release and Neurolysis 23	Totally restituted: 18 Improved: 3 Dead: 1 Did not respond: 1	
Freshwater, 1978		Number of patients with no symptoms at 2 years	Not significantly different by chi square test conducted by ECRI,
	Open Release 12	11	p = 0.64
	Release and Neurolysis 14	12	

^a: Two hands were lost to followup in the open release group and eleven in the neurolysis group. These hands were conservatively assumed to be unimproved. The significant loss to followup, as well as the fact that loss was not evenly distributed between groups, may render these data unreliable. This recalculation does not account for the additional 13 patients (14 hands) who were lost to followup for whom the group assignment was not reported.

b: Recalculated to account for patient attrition using the conservative assumption that treatment had failed for the two patients missing from the open release group and the one patient missing from the release and neurolysis group.

Table 72. Results of conservative meta-analysis of global outcome among patients treated with neurolysis for carpal tunnel syndrome

Author	Year	N	Effect Size	95% CI	p-value	Standardized Residual	Outlier by Std Residual?
Leinberry, 342	1997	50	0.089	-0.53-0.78	0.778	-0.64	No
Blair, 428	1996	75	0.067	-1.28-1.42	0.923	-0.30	No
Foulkes, 376	1994	36	0.432	-0.30-1.17	0.250	0.46	No
Mackinnon, ⁴⁰⁷	1991	63	0.282	-0.48-1.04	0.465	0.03	No
Lowry, ⁴²⁹	1988	50	0.140	-0.41-0.70	0.615	-0.52	No
Gelberman, 415	1987	61	0.697	0.11-1.28	0.019	1.61	No
Holmgren, ⁴¹⁶	1987	41	-0.741	-2.04-0.56	0.263	-1.56	No
Freshwater, 426	1978	26	0.324	-1.08-1.72	0.650	0.08	No
			Fixed effects model:				
			Overall Effect Size	95% CI	p-value of E.S.	Q	p-value of Q
			0.27	0.003-0.537	0.047	5.20	0.636

Figure 33. Results of meta-analysis of the effect of neurolysis on global outcome

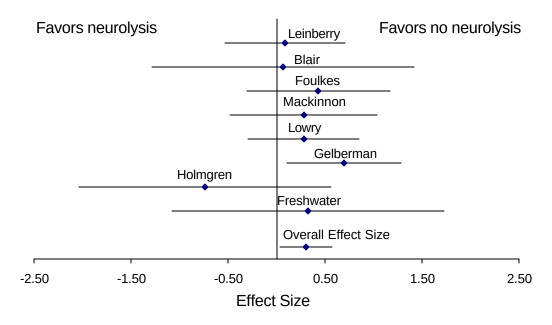


Table 73. Effect sizes of individual studies according to the assumptions used to calculate them

Study	Assumption used to calculate Hedges' d				
	Conservative No Recalculation		Anti- conservative		
Blair ⁴²⁸	0.067 (-1.28-1.42)	N/Aa	0.94 (-0.70-2.57)		
2.4	0.007 (-1.20-1.42)	IV/A ^a	` ′		
Foulkes 376	0.43 (-0.30-1.17)	0.30 (-1.53-2.13)	0.11 (-1.69-1.92)		
Lowry 429	0.14 (-0.41-0.70)	0.28 (-0.30-0.85)	0.37 (-0.19-0.93)		
Overall Effect Size	0.27 (0.003-0.537)	0.29 (0.01-0.97)b	0.31 (0.03-0.59)		
		0.28 (-0.01-0.57) ^c			

a: N/A; Not applicable. Data from this study were not recalculated.

Table 74. Effects of assumptions about individual studies on the overall effect size

	Study		Is the overall effect
Blair	Foulkes	Lowry	size significantly different from zero?
Conservative	Conservative	Conservative	Yes
Conservative	No Recalculation	No Recalculation	No
Conservative	Anti- conservative	Anti-conservative	No
Anti- conservative	Conservative	Conservative	Yes
Anti- conservative	No Recalculation	No Recalculation	Yes
Anti- conservative	Anti- conservative	Anti-conservative	Yes

Return to work

Two controlled trials, one of which was randomized, reported some information describing return to work. Both included patients who received bilateral procedures, and one had high (36%) attrition. Results are presented in Table 75 and summarized in Table 76. Neither study reported the number of patients who were working or on sick leave prior to treatment, so the number of patients returning to work could not be determined. As can be seen in Table 76 and Figure 35, both studies favor release without neurolysis, with the difference achieving statistical significance in one study. Because of incomplete reporting, no meta-analysis or power analysis was possible.

b: If the anticonservative effect size from the study by Blair is used.

c: If the conservative effect size from the study by Blair is used.

Table 75. Effect of neurolysis on return to work

Study	Number of Patients	Time to Return to Work	Statistical Significance of Difference Between Groups
Foulkes, 1994 376	Open Release	Median 53 Days (Range 1-180)	Not significantly different, stati stical test not reported.
	Release and Neurolysis	Median 59 Days (Range 14-120)	
	N not reported		
Freshwater, 1978 ⁴²⁶	N not reported	Stated only that patients receiving open release without neurolysis returned to work more quickly than those who received neurolysis.	This difference was statistically significant by the Mann-Whitney U test (p <0.01).

Table 76. Summary of the effect of neurolysis on return to work

Study	Which Procedure Had Faster Return to Work?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable)	Effect Size (95% Confidence Interval)
Foulkes, 1994 376	No neurolysis	No	Not calculable	Not calculable
Freshwater, 1978 426	No neurolysis	Yes	Not calculable	Not calculable

Table 77. Effect of neurolysis on carpal tunnel pain

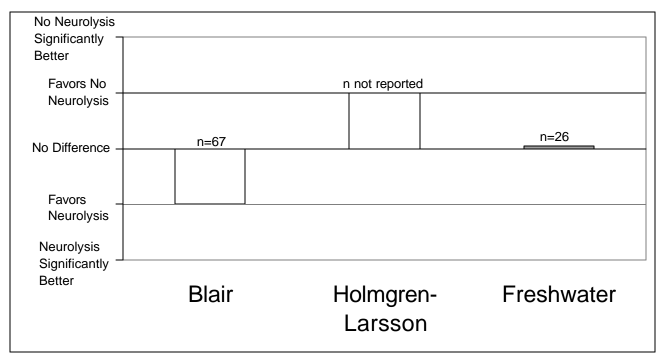
Study	Number of Hands	Pain	Statistical Significance of Difference Between Groups
Blair, 1996 428	Open Release 27	Preop: 25 had pain	Not significantly different by chi square
		Unimproved: 0 Improved: 8 (32%) No Pain: 17 (68%)	test conducted by ECRI, p = 0.11
		Preop: 42 had pain	
	Release and Neurolysis 48 (Hands)	Unimproved: 1 (2.4%) Improved: 5 (12%)	
		No Pain: 36 (86%)	
Holmgren- Larsson, 1985	48 Hands total; number in each group not reported.	Percent of patients reporting pain	Not reported
	Open release		
	Preop.	78	
	3-4 Weeks	0	
	6 Months	0	
	Release and neurolysis		
	Preop.	85	
	3-4 Weeks	4	
	6 Months	13	
Freshwater, 1978 ⁴²⁶		Patients with wrist pain:	Not significantly different by chi square test conducted by ECRI, p = 0.91
	Open Release 12	Preop: 2; Postop: 1	
	Release and Neurolysis 14	Preop 4; Postop: 1	Net classification of the control of
		Patients with night- waking pain and tenderness:	Not significantly different by chi square test conducted by ECRI, p = 0.97
	Open Release 12	Preop: 12; Postop: 0	
	Release and Neurolysis 14	Preop: 14; Postop: 0	

Table 78. Summary of the effect of neurolysis on pain

Study	Which Procedure led to less pain?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Blair, 1996 428	Neurolysis	No	28%	-0.57 (-1.23-0.10)
Holmgren- Larsson, 1985 433	No Neurolysis	No	Not calculable	Not calculable
Freshwater, 1978 ⁴²⁶	No difference	No	Not calculable	0.08 (-2.12-2.28)

a: Calculated by ECRI

Figure 36. Summary of effect of neurolysis on pain



An open bar indicates an RCT, a striped bar indicates a CT.

Table 79. Effect of neurolysis on hand function

Study	Number of Patients	Function	Statistical
			Significance of Difference Between
			Groups
Blair et al., 1996		Patients having difficulty:	There were no significant
420		Screwing Lids:	differences between groups before or after treatment
	Open Release 27	Preop: 25 (92.5%)	(test not reported)
		24 Months: 11 (40.7%)	
	Release and Neurolysis 48	Preop: 41 (85.4%)	
		24 Months: 15 (31.3%)	
		Picking up small objects:	
	Open Release 27	Preop: 18 (66.7%)	
		24 Months: 10 (37.0%)	
	Release and Neurolysis 48	Preop: 27 (56.3%)	
		24 Months: 9 (18.8%)	
		Lifting:	
	Open Release 27	Preop: 15 (55.6%)	
		24 Months: 7 (25.9%)	
	Release and Neurolysis 48	Preop: 25 (52.1%)	
	(Hands)	24 Months: 9 (18.8%)	
Foulkes et al.,	(Tanas)	Function rating (0-100)	Not reported
1994 ³⁷⁶	Open Release 8	Preop: 41	
	Open Neicase 0	29 Months: 89	
	10	Recalculated ^a : 79.4	
	Release and Neurolysis 15	Preop: 34	
		29 Months: 88	
8: Pocalculated by E	26	Recalculated: 65.2	1 4h - 1 - 1 1 1 1 1 1 1 1 1 1 1 - 1 1 1 1 1 1 1 1 1 1 1 - 1 1 1 1 1 1 1 1 1 1 1 - 1 1 1 1 1 1 1 1 1 1 1 - 1 1 1 1 1 1 1 1 1 1 1 - 1 1 1 1 1 1 1 1 1 1 1 - 1 1 1 1 1 1 1 1 1 1 1 - 1

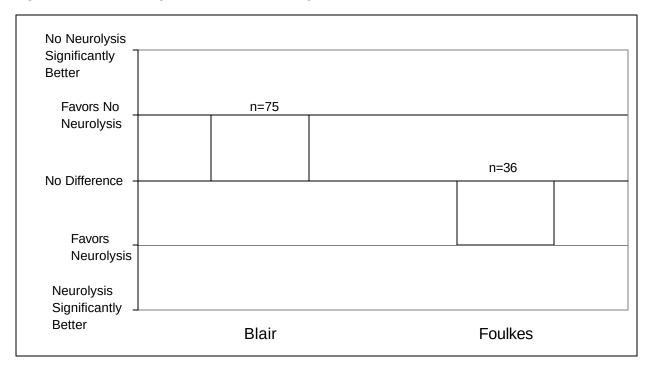
a: Recalculated by ECRI according to intent to treat principles by making the conservative assumption that the two patients lost to followup in the open release group had function ratings of 41 at 29 months, and the 11 lost to followup in the neurolysis group had function ratings of 34.

Table 80. Summary of effect of neurolysis on hand function

Study	Which Procedure led to superior function?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval)
Blair, 1996 ⁴²⁸	Neurolysis	No	Screwing Lids: 62% Picking up objects: 57% Lifting: 44%	Not calculable
Foulkes, 1994 376	Open release	Not reported	Not calculable	Not calculable

a: Calculated by ECRI

Figure 38. Summary of effect of neurolysis on hand function



Quality of Life

No studies reported on this outcome.

Harms

Only two randomized controlled trials reported on complications and adverse effects among patients receiving neurolysis. One of these had 50% attrition.³⁷⁶ These trials are listed below in Table 81. One controlled trial and one retrospective trial reported that there were no complications.^{415,426} There are insufficient data to allow one to reach an evidence-based conclusion.

Table 81. Complications in controlled trials of neurolysis for patients with carpal tunnel syndrome

Study	Group n	Complication	Number of patients reporting
Foulkes, 1994 ³⁷⁶	No Neurolysis 8 Hands	Infection	0
	Neurolysis 15 Hands	Infection	2
Lowry, 1988 ⁴²⁹	No Neurolysis 23	Persistent incisional pain Hand swelling Causalgia	3 0 1
	Neurolysis 24	Persistent incisional pain Hand swelling Causalgia	4 1 0

Conclusion

The available evidence suggest there is little or no benefit from performing neurolysis along with surgical release of the carpal tunnel. Meta-analysis of global outcomes demonstrates a benefit from not performing neurolysis that was not apparent from examination of the individual studies. Available return to work data also shows a trend toward an advantage of not performing neurolysis. There are insufficient data to allow one to reach an evidence-based conclusion, on the effect of neurolysis on pain or function. The possibility remains that neurolysis may be helpful is special cases, such as in the presence of marked scarring or neural adhesion, but no available evidence specifically documents the benefits and harms of neurolysis among such patients.

What are the relative benefits and harms of steroid injection into the carpal tunnel for persons with carpal tunnel syndrome?

Four prospective, randomized controlled trials describing 261 patients reported on the effect of steroid injections into the carpal tunnel.

Internal Validity

Three studies of steroid injections were double-blinded,^{36,452,453} and one was unblinded.⁴²⁷ Three studies assessed only one hand per patient, while Girlanda et al. assessed 53 hands in 32 patients.³⁶ This study therefore violated the statistical principle of independence between subjects. All four studies had no attrition and full compliance. Data on study internal validity may be found in Table 82.

Generalizability

None of the studies reported patient comorbidities, except when some comorbidities were excluded, as indicated by a zero in Table 83. Dammers, et al. excluded patients with mild disease. Results in this study may therefore be different from results in others. None of the studies provided information about patient employment characteristics.

Table 82. Internal validity of studies of steroid injection for carpal tunnel syndrome

Study	Number of patients	Percent of patients with bilateral CTS	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
O'Gradaigh, 2000 454	123	0%	Single	Not reported	RCT	No	0	Yes	100
Dammers, 1999 452	60	0%	Single	No	RCT	Double	0	Yes	100
Girlanda, 1993 ³⁶	32	65.6%	Single	Not reported	RCT	Double	0	Yes	100
Ozdogan, 1984 453	37	0%	Single	Not reported	RCT	Double	0	Yes	100

Table 83. Generalizability of studies of steroid injection for carpal tunnel syndrome

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
O'Gradaigh, 2000 ⁴⁵⁴	123	NR	NR	NR	NR	NR	0	NR	NR	NR	NR	No	No
Dammers, 1999 452	60	52	83.3	29	NR	NR	NR	NR	NR	NR	NR	No	Yes
Girlanda, 1993	32	45.5	81.3	53.5 (1-240)	0	0	NR	0	0	NR	0	No	No
Ozdogan, 1984 453	37	47.0	100	45.6	0	0	0	NR	NR	NR	0	No	No

Table 84. Effect of steroid injection on global outcome

Study	Number of Patients		Global Outcome	Statistical Significance of Difference Between Groups			
O'Gradaigh, 2000 454			Patients showing improvement of symptoms	Treatments were superior to controls at either time point by chi square test, p <0.05			
			6 Weeks:	Treatments were not significantly different from each other at either			
	No Injection 20 mg Triamcinolone 25 mg Hydrocortisone 100 mg Hydrocortisone	20 18 32 53	1 (5.0%) 13 (72.2%) 21 (65.6%) 34 (64.1%)	time point by chi square test, p >0.05.			
			6 Months:				
	No Injection 20 mg Triamcinolone 25 mg Hydrocortisone 100 mg Hydrocortisone	20 18 32 53	0 (0%) 8 (44.4%) 14 (43.8%) 17 (32.1%)				
Dammers, 1999 ⁴⁵²			Patients with No symptoms or minor symptoms	Treatments were significantly different at both time points (p = 0.000011 and 0.0002 respectively, chi square test conducted by			
			1 Month	ECRI)			
	Placebo (10 mg Lignocaine)	30	6 (20.0%)				
	10 mg Lignocaine and 40 mg Methylprednisone	30	23 (76.7%) 12 Months				
	Placebo (10 mg Lignocaine)	30	2 (6.7%)				
	10 mg Lignocaine and 40 mg Methylprednisone	30	15 (50.0%)				
Girlanda, et al., 1993 ³⁶	Wettrypredmisone		Mean symptom score (0-10)	Not reported			
			Pretreatment:				
	Placebo (Saline)	26	9				
	15 mg Methylprednisone	27	8				
	wietriyipi euriisorie		1 Week				
	Placebo (Saline)	26	7				

Study	Number of Patient	:s	Global Outcome	Statistical Significance of Difference Between Groups
	15 mg Methylprednisone	27	3 2 Months	
	Placebo (Saline)	26	8	
	15 mg Methylprednisone	27	1.5	
Ozdogan and Yazici, 1984 ⁴⁵³	1.5mg Betamethasone in the deltoid muscle	19	Pretreatment:: Severe 13 Moderate 4 Minimal 2 No Symptoms 0	Groups were not significantly different, p = 0.83, chi square test conducted by ECRI
	1.5mg Betamethasone in the carpal tunnel	18	Severe 11 Moderate 6 Minimal 1 No Symptoms 0	
	1.5mg Betamethasone in the deltoid muscle	19	1 Week: Severe 5 Moderate 2 Minimal 8 No Symptoms 4	Groups were not significantly different, p = 0.25, chi square test conducted by ECRI.
	1.5mg Betamethasone in the carpal tunnel	18	Severe 2 Moderate 3 Minimal 8 No Symptoms 5	
	1.5mg Betamethasone in the deltoid muscle	19	1 Month: Severe 8 Moderate 8 Minimal 2 No Symptoms 1	Groups were significantly different, p = 0.009, chi square test conducted by ECRI
	1.5mg Betamethasone in the carpal tunnel 18	18	Severe 6 Moderate 3 Minimal 0 No Symptoms 9	

Table 85. Summary of effect of steroid injection on global outcome

Study	Which Procedure led to Superior Global Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
O'Gradaigh, 2000 ⁴⁵⁴	Injection	Yes	20 mg Triamcinolone 22%	20 mg Triamcinolone 6 Weeks: 2.11 (0.86 – 3.35) 6 Months: 1.89 (0.27 – 3.52)
			25 mg Hydrocortisone 18%	25 mg Hydrocortisone 6 Weeks: 1.95 (0.77 - 3.13) 6 Months: 1.88 (0.29 – 3.48)
			100 mg Hydrocortisone 17%	100 mg Hydrocortisone 6 Weeks: 1.92 (0.77 – 3.07) 6 Months: 1.62 (0.05 – 3.20)
Dammers, 1999 452	Injection	Yes	16%	1 Month: 1.40 (0.720-02.08) 12 Months: 1.44 (0.55 – 2.32)
Girlanda, 1993 ³⁶	Injection	Not reported	Not calculable	Not calculable
Ozdogan, 1984 ⁴⁵³	Injection	At 1 month only	Not calculable	1 Week: 0.25 (-0.39 – 0.90) 1 Month: 0.28 (-0.37 – 0.40)

a: Calculated by ECRI

Table 86. Internal validity of studies of oral medication for carpal tunnel syndrome

Study	Number of patients	Percent of patients with bilateral CTS	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
Chang, 1998 35	91	0%	Single	Not reported	RCT	Double	18	No	NR
Herskovitz, 1995 ⁴⁵⁵	18	0%	Single	Not reported	RCT	Double	3	No	NR

Table 87. Generalizability of studies of oral medication for carpal tunnel syndrome

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Chang, 1998 35	91	45.7	58.2	NR	0	0	NR	NR	0	0	NR	Yes	Yes
Herskovitz, 1995	18	49.6	80.0	20.6	6.6	6.6	NR	0	0	NR	NR	Yes	Yes

Table 88. Oral drugs used to treat carpal tunnel syndrome in controlled studies

Drug	Dose	Description
Prednisone	20mg/day for 1 week, then 10mg/day for 1 week	An anti-inflammatory steroid
Prednisolone	20mg/day for 2 weeks, then 10mg/day for 2 weeks	An anti-inflammatory steroid
Tenoxicam	20mg/day for 4 weeks	A nonsteroidal anti-inflammatory drug (NSAID)
Trichlormethiazide	2mg/day for 4 weeks	A diuretic, used to reduce swelling and lower carpal tunnel pressure

Global outcome

Both studies reported global symptom scores. This was the mean of five symptom severity ratings on a scale of zero to ten. The symptoms rated were pain, numbness, paresthesia, weakness/clumsiness and nocturnal awakening. These data are summarized in Table 89. As can be seen in Table 90 and Figure 40, both reports found statistically significant decreases in symptom scores among patients treated with steroids compared to placebo controls. However, Herskovitz et al. reported that symptoms returned after the cessation of treatment. In neither study did symptom scores approach zero, indicating that although there was some relief, symptoms were still present. Chang et al. reported a 64% mean decrease in global symptom scores, while Herskovitz et al reported a 68% decrease. Neither paper indicated whether the patients were satisfied with their level of symptom relief.

When the data were recalculated to account for patient attrition, the steroid groups in both studies still showed a greater than 50% reduction in global symptom scores. However, because we are unable to accurately estimate the standard deviations around the recalculated means, we are unable to determine whether the difference remains statistically significant. The number of patients reporting symptom relief in the report by Herskovitz is not statistically significantly different between groups once we attempted to compensate for patient attrition by assuming that patients for whom there was no data did not improve.

In the study by Chang, neither the diuretic nor the NSAID caused statistically significant symptom relief compared to placebo control. However, a single small trial with high loss to followup is not sufficient proof that these agents have no effect. Moreover, only a single dosage of each drug was tested. There are no published data on the effectiveness of these agents at other dosages. The power of the study by Chang was sufficient to detect medium-sized (20-30%) differences between groups. The differences between placebo and steroid were greater than this, while the differences between the other groups and placebo were too small to be statistically significant with the available power.

Table 89. Effect of oral medications on global outcome of carpal tunnel syndrome

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Chang, et al., 1998 35		Mean global symptom score ^a	
	Placebo 16	Baseline: 22.9±5.9 2 Weeks: 21.6±6.4 4 Weeks: 20.8±6.6	Symptom reduction among patients receiving steroid was significantly greater at 2 weeks than among patients in the other three groups (F = 7.37, p = 0.0002) Symptom reduction among patients
	Diuretic 16	Baseline: 26.0±3.8 2 Weeks: 22.3±5.5 4 Weeks: 21.6±6.3	receiving steroid was significantly greater at 4 weeks than among patients in the other three groups (F = 10.7, p = 0.0001) NSAID and diuretic groups were not significantly different from placebo at either time point.
	NSAID 18	Baseline: 29.7±8.4 2 Weeks: 24.7±8.6 4 Weeks: 24.0±9.7	
	Steroid 23 (Prednisolone)	Baseline: 27.9±6.9 2 Weeks: 15.0±6.8 4 Weeks: 10.0±7.5	
	Recalculated ^b Placebo 23	Recalculated ^b Baseline: 22.9 2 Weeks: 22.0 4 Weeks: 21.4	
Harakavitz et al. 1005 (55	Steroid 26	Baseline: 27.9 2 Weeks: 16.5 4 Weeks: 12.1	
Herskovitz, et al., 1995 ⁴⁵⁵	Placebo 9	Mean global symptom score ^a Baseline: 23 2 Weeks: 19 4 Weeks: 17 8 Weeks: 16.5	Groups were significantly different only at 2 weeks (p <0.05, t test)

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
	Steroid 6 (Prednisone)	Baseline: 25 2 Weeks: 8 4 Weeks: 11 8 Weeks: 20	
	Recalculated ^b	Recalculated ^b	
	Placebo 10	Baseline: 23 2 Weeks: 19.4 4 Weeks: 17.6 8 Weeks: 17.2	
	Steroid 8	Baseline: 25 2 Weeks: 12.3 4 Weeks: 14.5 8 Weeks: 21.3	
		Number of patients reporting improvement in symptoms:	Numbers were the same for all time points, and were significantly different between groups (p = 0.02, test not reported)
	Placebo 9	3	Improvement rates were no longer statistically significant if the two patients
	Steroid 6 (Prednisone)	6	from the steroid group and one from the placebo group who were not reported on were assumed not to have improved, p = 0.058 by chi square test conducted by ECRI.

a: The sum of severity ratings (scale 0-10) for 5 symptoms: pain, numbness, paresthesia, weakness/clumsines s, and nocturnal wakening

b: Recalculated to account for patient attrition using the conservative assumption that patients for whom no data was provided had scores equal to the mean baseline score for that group.

Table 90. Summary of effect of oral medications on global outcome of carpal tunnel syndrome

Study	Which Medication led to Superior Global Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Chang,	Steroid	Yes	Diuretic	Diuretic
1998 ³⁵			2 Weeks: 20.0%	2 Weeks: -0.11 (-0.81 – 0.58)
			4 Weeks: 22.4%	4 Weeks: -0.12 (-0.81 – 0.57)
			NSAID	NSAID
			2 Weeks: 24.6%	2 Weeks: -0.40 (-1.08 – 0.28)
			4 Weeks: 27.9%	4 Weeks: -0.37 (-1.05 – 0.31)
			Steroid	Steroid
			2 Weeks: 20.2%	2 Weeks: 0.97 (0.30 – 1.65)
			4 Weeks: 22.4%	4 Weeks: 1.48 (0.76 – 2.20)
Herskovitz,	Steroid	Yes	Global Symptom Score	Global Symptom Score
1995 ⁴⁵⁵			Not calculable	2 Weeks: 1.08 (-0.03 – 2.18)b
			Number of Patients	Number of Patients Improved
			Improved 49%	1.65 (-0.09 – 3.39)

a: Calculated by ECRI
b: Estimated by ECRI based on the conservative assumption that p = 0.049.

Quality of Life

Neither study reported this outcome.

Harms

Chang et al. reported the number of patients experiencing nausea and epigastric pain, while Herskovitz et al. reported the number experiencing any perceived effect. These results are presented in Table 91. In both studies, numbers of patients reporting side effects were not significantly different between treated groups and placebo groups by chi square test conducted by ECRI (p > 0.3). However, there are too few studies to allow one to reach a firm evidence-based conclusion about the side effects experienced by patients with carpal tunnel syndrome who are given oral medications.

Table 91. Side effects of oral medications for carpal tunnel syndrome

Study	Group	Complication	Number of patients experiencing complication
Chang, et al., 1998 35	Placebo 16	Nausea Epigastric pain	1 2
	Diuretic 16	Nausea Epigastric pain	0 2
	NSAID 18	Nausea Epigastric pain	3 3
	Steroid 23 (Prednisolone)	Nausea Epigastric pain	3 2
Herskovitz, et al., 1995 ⁴⁵⁵	Placebo 9	Nausea/abdominal discomfort, constipation, insomnia, headache, dysuria, and burning nostrils	3
	Prednisone 6	Nausea/abdominal discomfort, constipation, dysgeusia, mild hypoglycemia	3

Conclusions

Two double-blinded randomized controlled trials suggest that oral steroids may lead to a reduction in symptoms of CTS. A single published randomized controlled trial indicates that oral tenoxicam and trichlormethiazide do not reduce the symptoms of CTS under the dosing regimens described. The effects of oral steroids are short-lived and may not be sufficient for patient satisfaction. There are no published controlled trials describing the effects of higher doses or longer treatment regimens.

Table 92. Internal validity of the study comparing oral and injected steroids

Study	Number of patients	Percent bilateral patients	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
Wong, 2001	60	23.3%	Single	Not	RCT	Double	0	Yes	NR
456				Reported					

Table 93. Generalizability of the study comparing oral and injected steroids for carpal tunnel syndrome

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Wong, 2001 456	60	49	88.3%	NR	0	0	0	0	0	0	0	Yes	Yes

Global Outcome

The outcome measure was global symptom score, the sum of ratings (0 to 10) of pain, numbness, paresthesia, weakness/clumsiness and nocturnal awakening. These scores are given in Table 94, and the results are summarized in Table 95. This outcome was statistically significantly different between groups at 8 weeks and 12 weeks. The difference between groups at two weeks was smaller than the study had the power to detect.

Table 94. Relative effect of steroid injection and oral steroids on global outcome of CTS

Study	Number of Patients	Global Symptom Score	Statistical Significance of Difference Between Groups
Wong, 2001 456	Injection 30		Groups were significantly different at 8 weeks and 12 weeks by t-test conducted by ECRI.
	Pretreatment	25.00±6.41	and 12 weeks by Flest Conducted by LCIVI.
	2 Weeks	13.57±7.47	
	8 Weeks	13.67±8.27	
	12 Weeks	14.30±8.42	
	Oral 30		
	Pretreatment	25.73±8.31	p = 0.705
	2 Weeks	17.77±9.98	p = 0.070
	8 Weeks	20.83±8.73	p = 0.0019
	12 Weeks	21.40±9.64	p = 0.0036

Table 95. Summary of the relative effect of steroid injection and oral steroids on global outcome of CTS

Study	Which Procedure led to Superior Global Outcome?	Was the Difference Statiscally Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Wong, 2001	2 Weeks: Injection	No	21%	0.47 (-0.09-1.03)
	8 Weeks: Injection	Yes	21%	0.831. (0.25-1.41)
	12 Weeks: Injection	Yes	22%	0.77 (0.20-1.35)

a: Calculated by ECRI

Return to Work

This study did not report this outcome.

Return to Activities of Daily Living

This study did not report this outcome.

Pain

This study did not report this outcome.

Function

This study did not report this outcome.

Quality of Life

This study did not report this outcome.

<u>Harms</u>

Harms reported among the two groups are given in Table 96. Steroid and placebo injection led to injection pain in two patients each. All other side effects were reported to have been experienced by the oral steroid group only. The difference in occurrence of side effects between groups was statistically significant by chi square test conducted by ECRI (p = 0.0195).

Table 96. Reported harms of injected and oral steroids

Study	Group	Complication	Number of patients experiencing complication
Wong, 2001 456	Injected 30	Injection pain Increased appetite	0
		Bloating	0
		Insomnia	0
	Oral 30	Injection pain	2
		Increased appetite	3
		Bloating	2
		Insomnia	2

Conclusions

Although only a single study, this study had high internal validity, providing evidence that, under the conditions of the experiment, steroid injection leads to greater reduction of symptoms with fewer side effects than oral steroid. The experiment is short-term (12 weeks) and does not address the issue of whether the effect of injection remains effective at longer time points. Further, it does not address whether continued treatment with oral steroids leads to further benefits or harms to the patient.

What are the relative benefits and harms of physical therapy for persons with carpal tunnel syndrome?

Two randomized controlled trials describing 121 patients reported on the effects of various forms of physical therapy. Tal-Akabi and Rushton compared groups receiving nerve mobilization, groups receiving bone mobilization and a no-treatment control group. Provinciali et al. compared a program of physical therapy including strengthening exercises, massage, gliding exercises and sensory re-training to instruction in a program of home-based strengthening exercises.

Internal Validity

The study by Provinciali was rater-blinded, while the other was unblinded. Trial characteristics affecting internal validity are listed in Table 97. Neither study had any reported attrition, and neither reported on patient compliance.

Generalizeability

In both studies, patients were predominantly middle-aged (mean 54.8 years) and female (67%-82%), as reported in Table 98. This is consistent with the overall population with CTS as described in the introduction under Epidemiology. Tal-Akabi excluded patients with comorbidities, while Provincialli did not report comorbidities. Both studies excluded patients with mild disease. This may limit generalizability, as patients with mild disease are more likely to receive noninvasive treatments such as physical therapy than patients with severe disease, who may be candidates for surgery. Neither study reported patient employment characteristics.

Table 97. Internal validity of studies of physical therapy for carpal tunnel syndrome

Study	Number of patients	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
Provinciali, 2000 ⁴²⁷	100	Single	Not reported	RCT	Rater	0	Yes	NR
Tal-Akabi, 2000 ⁴⁵⁷	21	Not Reported	Not reported	RCT	No	0	Yes	NR

Table 98. Generalizability of studies of physical therapy for carpal tunnel syndrome

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Provincialli, 2000 ⁴²⁷	100	56.45 (24-86)	82.0	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes
Tal-Akabi, 2000	21	47.1 (29-85)	66.6	27.6 (12-36)	0	0	0	0	0	0	NR	No	Yes

Table 99. Global outcome of physical therapy for carpal tunnel syndrome

Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Tal-Akabi and Rushton, 2000 ⁴⁵⁷	Neurodynamic mobilization 7	Global Score (Number of patients going on to receive surgery)	The two treated groups were not significantly different from each other (p = 0.51 by chi square test conducted by ECRI); both were significantly different from control (p = 0.03 and 0.008, respectively).
	Carpal Bone mobilization 7	1	
	No treatment (Control) 7	6	

Table 100. Summary of Global outcome of physical therapy for carpal tunnel syndrome

Study	Which Procedure led to Superior Global Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Tal-Akabi, 2000 ⁴⁵⁷	Carpal bone mobilization	Yes	50%	Neurodynamic mobilization 1.40 (-0.08 – 2.87) Carpal bone mobilization 1.85 (0.20 – 3.50) Difference between-treatment groups 0.45 (-1.42-1.93)

a: Calculated by ECRI

Return to work

A single study reported time to return to work. Provincialli et al. reported that patients receiving physical therapy returned to work earlier than patients assigned to home exercise. As can be seen in Table 101, the difference was statistically significant, but the number of patients for whom this measurement was taken was not reported. Further, it is unclear exactly what was measured. These numbers are described both as time to return to daily activities and time to return to work. These ambiguities render it difficult to draw conclusions from these data.

Table 101. Time to return to work after physical therapy for carpal tunnel syndrome

Study	Number of Patients	Days until Return to Activities of Daily Living	Statistical Significance of Difference Between Groups
Provincialli et al., 2000 ⁴²⁷	Physical Therapy Home Exercise	32.16±10.72 42.55±13.39	Difference was statistically significant by ANOVA (p <0.006)
	Number of patients is unknown because patients receiving workers' compensation were excluded. The number of such patients was not reported.		

Return to Activities of Daily Living

This outcome was not reported by either study.

Pain

Both studies reported pain scores. Tal-Akabi and Rushton also reported pain relief scores. These data are given in Table 102. Provincialli et al. found no statistically significant difference between the program of physical therapy and home exercise instructions. Tal-Akabi and Rushton found that one treatment, carpal bone mobilization, but not the other treatment, neurodynamic modulation, led to pain scores statistically significantly lower than those in the control group (p = 0.003 and 0.35 respectively). The two treatment groups were not significantly different from each other (p = 0.18). The study lacked the statistical power to detect the difference between these groups. Only large between group differences (>50%) could be detected in this study, as can be seen in Table 103. While the differences between carpal bone mobilization and control are large enough to be detected, other between group differences are not. The fact that carpal bone mobilization led to a statistically significant effect while neurodynamic mobilization did not suggests, but does not prove, that carpal bone manipulation is the superior treatment for pain. Further study is necessary to test the differences between these therapies.

Although pain ratings in the VAS group were not significantly different from control after treatment, differences between pain relief scores were statistically significant. It is unclear which is the superior measure of pain.

Table 102. Effects of nerve and bone mobilization on pain from carpal tunnel syndrome

Study	Number of Patients	Pain	Statistical Significance of Difference Between Groups
Provinciali, 2000	Physical	Sum of patients' pain ratings (scale	Groups were not significantly different by chi square test (p >0.001; p-level required
	Therapy 50	not reported)	for significance adjusted by Provinciali using the Bonferroni correction related to
	Pretreatment	149	40 comparisons)
	1 Month	55	
	2 Months	50	
	Home Exercise 50		
	Pretreatment	145	
	1 Month	54	
	2 Months	50	
Tal-Akabi, 2000	Neurodynamic mobilization 7	Pain (VAS, 0-10) Baseline 2.42±1.51 3 Weeks 1.57±1.4	After treatment, the carpal bone mobilization group was significantly different from control by t-test conducted by ECRI (p = 0.003), but the neurodynamic mobilization group was not significantly different from contol (p = 0.35) or from carpal bone mobilization (p = 0.18)
	Carpal Bone mobilization 7	Baseline 2.29±0.95 3 Weeks 0.71±0.76	
	No treatment (Control) 7	Baseline 2.0±1.29 3 Weeks 2.14±0.69	
	Neurodynamic mobilization 7 Carpal Bone mobilization 7	Pain Relief Rating 3.14±1.35 3.71±0.95	Not significantly different between the two treated groups (p = 0.38), but both the neurodynamic mobilization group and the carpal bone mobilization group were significantly different from control (p = 0.00005 and 0.0000002, respectively)
	No treatment	0±0	

Study	Number of Patients	Pain	Statistical Significance of Difference Between Groups
	(Control) 7		

Table 103. Summary of effects of nerve and bone mobilization on pain from carpal tunnel syndrome

Study	Which Procedure led to Less Pain?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Provinciali, 2000 ⁴²⁷	No difference	No	Not calculable	Not calculable
Tal-Akabi, 2000 ⁴⁵⁷	Carpal bone mobilization	Yes	Neurodynamic mobilization: 60% Carpal bone mobilization: 54%	VAS Neurodynamic mobilization 0.48 (-0.62 – 1.58) Carpal bone mobilization 1.84 (0.59 – 3.10) Pain Relief Rating Neurodynamic mobilization 3.08 (1.53 – 4.63)
				Carpal bone mobilization 5.17 (2.99 – 7.35)

a: Calculated by ECRI

Function

In the study by Provincialli, function was measured using a nine-hole peg test. Function scores were not significantly different between groups at any time point. In the study by Tal-Akabi and Rushton, functional scores were based on the impairment rating of the patient's most impaired activity. Thus, a lower score indicates superior function. These scores were not significantly different before treatment. Results are presented in Table 104. After treatment, functional scores in the carpal bone mobilization group were significantly lower than those of the control group (p = 0.01), while those of the neurodynamic mobilization group were not (p = 0.09). The two treatment groups were not significantly different from each other (p = 0.57). As presented in Table 105, the study only had the power to detect large (>50%) differences between groups. Only the difference between carpal bone mobilization and control was large enough to be found statistically significant.

Table 104. Effect of physical therapy on function

Study	Number of Patients	Function	Statistical Significance of Difference Between Groups
Provinciali et al. 2000, ⁴²⁷	Physical Therapy 50 Pretreatment 12 Days 1 Month 2 Months Home Exercise 50 Pretreatment 12 Days 1 Month	Time (units not stated) to complete nine-hole peg test 22.35±5.14 23.8a 20.5 19.5 22.38±3.23 20.5 19	Groups were not significantly different by t test (p >0.001; p-level required for significance adjusted by Provinciali using the Bonnferoni correction related to 40 comparisons
Tal-Akabi and Rushton, 2000 ⁴⁵⁷	2 Months Neurodynamic mobilization 7 Carpal Bone mobilization 7 No treatment (Control) 7	Function Score (Range 0-4) Baseline 2.0±1.41 3 Weeks 1.14±1.35 Baseline 2.0±1.41 3 Weeks 0.71±0.76 Baseline 2.42±1.27 3 Weeks 2.42±1.27	After treatment, carpal bone mobilization group was significantly different from control group (p = 0.01) neurodynamic mobilization group was not (p = 0.09). The two treatment groups were not significantly different from each other (p = 0.57). Etests conducted by ECRI.

a: Estimated by ECRI from a published chart

Table 105. Summary of the effect of physical therapy on function

Study	Which Procedure led to Superior Function?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Provinciali, 2000 ⁴²⁷	No difference	No	Not calculable	Not calculable
Tal-Akabi, 2000 ⁴⁵⁷	Carpal bone mobilization	Yes	Neurodynamic mobilization 63%	Neurodynamic mobilization 0.91 (-0.21 – 2.19)
			Carpal bone mobilization 50%	Carpal bone mobilization 1.53 (0.34 – 2.72)

a: Calculated by ECRI

Quality of Life

This outcome was not reported by either study.

Harms

No harms were reported by either study.

Conclusions

Manual therapy may have some use in the treatment of carpal tunnel syndrome. A single study suggests that carpal bone mobilization provides pain relief, improves function, and delays or eliminates the need for surgery among patients with carpal tunnel syndrome. As Results from neurodynamic mobilization show a similar trend, but because of a lack of statistical power one cannot conclude that this trend is real. For the same reason, differences in effectiveness between these two treatment groups cannot be determined. The study was not placebo-controlled and was not blinded. The observed effects may have been influenced by a placebo effect or rater bias.

A larger, more statistically powerful study found no difference between the effects of a physical therapy program and home exercise instructions on pain or function. However, patients receiving physical therapy returned to work faster than those instructed to exercise at home.

Although these studies indicate a trend toward physical therapy having an effect on carpal tunnel syndrome, they are too small and inconclusive for one to reach a firm evidence-based conclusion.

What are the relative benefits and harms of ultrasound for persons with carpal tunnel syndrome?

One patient-blinded randomized controlled trial describing 18 patients reported on the effects of ultrasound.³³ This study compared two different levels of intensity of ultrasound to placebo.

Internal Validity

Factors affecting the internal validity of this study are listed in Table 106. The data are reported in terms of the number of hands, rather than number of patients, and among the 18 patients, 30 hands were treated. This violates statistical assumptions of independence.

Generalizability

As can be seen in Table 107, the 18 patients were middle-aged (range 37-66), and all were female. Patients with comorbidities were excluded, as were patients with very mild or severe CTS. These exclusions may limit the generalizability of the trial's results, especially given the fact that only a single trial has been published.

Table 106. Internal validity of the study of ultrasound for carpal tunnel syndrome

Study	Number of patients	Percent bilateral patients	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
Oztas, 1998 33	18	66.7%	Single	No	RCT	Patient	0	Yes	NR

Table 107. Generalizability of the study of ultrasound for carpal tunnel syndrome

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Oztas, 1998 33	18	51.6 (37-66)	100	84 (6-240)	0	0	0	0	0	0	0	Yes	Yes

Table 108. Effects of ultrasound on carpal tunnel syndrome

Study	Number of Hands ^a	Outcome	Statistical Significance of Difference Between Groups
Oztas, et al., 1998 33		Pain (VAS, 0-10)	
	1.5 W/cm² 10	Baseline 6.10±2.50 Posttreatmen® 2.90±1.69	All posttreatment scores were significantly different from baseline (p <0.05, t test). There were no significant differences between groups (p >0.05, 1-way ANOVA).
	0.8 W/cm² 10	Baseline 7.10±2.38 Posttreatment 3.60±1.90	
	0 W/cm² (Placebo) 10	Baseline 7.90±1.80 Posttreatment 4.00±2.40	
		Global Outcome (Mean of a categorical symptom rating, 0-3 scale)	
	1.5 W/cm² 10	Baseline 2.30±0.68 Posttreatment 1.40±0.52	
	0.8 W/cm² 10	Baseline 2.60±0.70 Posttreatment 1.70±0.82	
	0 W/cm² (Placebo) 10	Baseline 2.60±0.69 Posttreatment 1.40±0.97	

a: Eighteen patients with a total of 30 affected hands were treated.
 b: Followup time was five days after two weeks of treatment

Table 109. Summary of effects of ultrasound on carpal tunnel syndrome

Study	Which Procedure led to Superior Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Oztas, 1998 33	No differences	No	Pain 49%	Pain 1.5 W/cm² 0.51 (-0.38 – 1.40) 0.8 W/cm² 0.18 (-0.70 – 1.06)
			Global Outcome 52%	Global Outcome 1.5 W/cm² 0 (-0.88 – 0.88) 0.8 W/cm² -0.32 (-1.20 – 0.56)

a: Calculated by ECRI

Conclusions

Only one study meeting inclusion criteria addresses the use of ultrasound for carpal tunnel syndrome. Because of this, and because its design and analysis difficulties, one cannot reach a firm evidence-based conclusion.

What are the relative benefits and harms of full-time and nighttime-only splint use for persons with carpal tunnel syndrome?

A single unblinded randomized trial of 21 patients compared the effects of nighttime-only and full-time splint use.³⁴

Internal Validity

Study characteristics related to internal validity are presented in Table 110. This study reported a 20% loss to followup. Of those patients who returned for followup, there was considerable noncompliance. Only 85% of the nighttime-only group reported complete or nearly complete nighttime splint use. Twenty-three percent of this group also reported some daytime use, despite instructions to wear the splint only at night. Complete or nearly-complete daytime use was reported by only 27% of patients instructed to wear the splints full-time. Nearly 43% of the patients had bilateral CTS, and results were reported per hand rather than per patient. This, combined with the loss to followup and noncompliance issues, raises serious doubts as to the reliability of the results of this study.

Generalizability

Patients were middle age (mean 60 years) and predominantly male. This distinguishes them from the majority of CTS patients, who are usually female. Patient characteristics are listed in Table 111. No information about comorbidities or employment characteristics was reported, except that 57.1% of patients were employed (Table 112).

Table 110. Internal validity of the study of full-time and nighttime-only splint use for carpal tunnel syndrome

Study	Number of patients	Percent of bilateral patients	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
Walker, 2000 ³⁴	21	42.9%	Single	Not reported	RCT	No	4	No	14

Table 111. Generalizability of the study of full-time and nighttime-only splint use for carpal tunnel syndrome

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Walker, 2000 34	21	60	3.0	28.5	NR	NR	NR	NR	NR	NR	NR	No	Yes

Table 112. Patient employment characteristics in the study of full-time and nighttime-only splint use for carpal tunnel syndrome

Study	Number of patients	% Patients employed	% Patients receiving workers' compensation	% Patients retired	% Patients Homemakers	Reported
Walker, 200	0 34 21	57.1	Not reported	Not reported	Not reported	Not reported

Because there is only a single study reporting two outcomes, we discuss the results together. No results were described for return to work, return to ADLs, pain, quality of life or harms. Reported results can be found in Table 113. There were no statistically significant differences between groups in global outcome or functional ability, as can be seen in Table 114. However, the study lacked the statistical power to detect small differences between groups. Only medium (28%-33%) or larger differences would have been statistically significant.

Table 113. Results of comparison between full-time and part-time splint wear for carpal tunnel syndrome

Study	Number of Hands	Outcome	Statistical Significance of Difference Between Groups
Walker et		Global outcome	Change from pre to post was not significantly
al., 2000 34		(Symptom	different between groups by t-test. p-values were
		severity)	not reported.
	Nighttime-only 13		
	Pretest	2.89±0.96	
	Posttest	2.30±0.93	
	Full-time 11		
	Pretest	2.79±0.69	
	Posttest	2.09±0.62	
	1 datioat	2.07±0.02	
		Functional	Change from pre to post was not significantly
		(Levine) score	different between groups by t-test. p-values were
	Nighttime-only 13		not reported.
	Pretest	2.75±1.01	
	Posttest	2.14±0.87	
	Full-time 11		
	Pretest	2.27±1.03	
	Posttest	1.93±0.77	

Table 114. Summary of comparison between full-time and part-time splint wear for carpal tunnel syndrome

Study	Which Procedure led to Superior Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a		
Walker et al., 2000 ³⁴	Full-time use	No	Global outcome 29%	Global outcome 0.25 (-0.55 – 1.06)		
			Functional (Levine) score 33%	Functional (Levine) score 0.25 (-0.56 – 1.05)		

a: Calculated by ECRI

Conclusions

Splint use was addressed only by a single trial that had design difficulties. Because of this, one cannot reach an evidence-based conclusion about splint use.

What are the relative benefits and harms of open carpal tunnel release with ligament reconstruction for persons with carpal tunnel syndrome?

One non-blinded, retrospective controlled trial reported on the effects of ligament lengthening or reconstruction. ⁴⁸

Internal Validity

The study did not include patients with bilateral CTS, meaning that there were no violations of the assumption of statistical independence. There was no attrition. Therefore intent-to-treat principles were followed. Study characteristics related to internal validity are listed in Table 115.

Generalizability

Patients were predominantly female and the reported range of ages (24-88 years) is broadly similar to that of the overall CTS population. The trial did not describe patient comorbidities or employment characteristics.⁴⁸ Patient characteristics are presented in Table 116.

Table 115. Internal validity of studies of open carpal tunnel release with and without ligament reconstruction

Study	Number of patients	Percent of bilateral patients	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?
Karlsson, 1997 ⁴⁸	74	0%	Single	Not reported	Retro	No	0	Yes

Table 116. Generalizability of studies of open carpal tunnel release with and without ligament reconstruction

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Karlsson, 1997 48	74	NR; (24-88)	59.6	[Median) 6 (1-60)	NR	NR	NR	NR	NR	NR	NR	No	No

Time to return to work among patients treated with open release or ligament reconstruction is reported in Table 117. No other patient-oriented outcomes were reported.

Patients who received ligament reconstruction were statistically significantly slower to return to work than those who received open release without ligament reconstruction. The effect size was statistically significantly different from zero (d = 0.65, 95% C.I. = 0.15 - 1.15).

Table 117. Effect of ligament reconstruction on time to return to work

Study	Number of Patients	Weeks until Return to Work	Statistical Significance of Difference Between Groups
Karlsson et al., 1997 48	Open release 50 Release and reconstruction 24	4.5 (Range 1-12) 6.0 (Range 3-24)	Groups were significantly different (p <0.01, t-test.).

Conclusions

The results of one study suggest that suboptimal outcomes are obtained when patients receive ligament reconstruction. However, this trial was neither randomized nor blinded, so one cannot draw firm evidence-based conclusions from it.

What are the relative benefits and harms of open carpal tunnel release with early or late mobilization for persons with carpal tunnel syndrome?

Three prospective, randomized controlled trials describing 171 patients compared early and late mobilization (removal of cast or splint) after open carpal tunnel release.

Internal Validity

None of these trials were blinded. Study characteristics related to internal validity are presented in Table 118. Only one study had patient attrition, and two reported results of bilateral patients as per hand rather than per patient. One study had a high (92.7%) rate of compliance, while the other two did not report compliance.

Generalizability

Patient characteristics are reported in Table 119. The studies by Finsen and Bury included predominantly female, middle-aged patients, while Cook did not report these characteristics. The studies differed in their inclusion/exclusion criteria, with Bury et al excluding patients with mild carpal tunnel syndrome, ⁴⁵⁸ Cook et al. excluding both the most mild and the most severe cases, ⁴³² and Finsen et al. not excluding according to severity. ³¹⁹ Finsen and Cook excluded patients with comorbidities, while Bury included patients with other nerve impingement conditions. These differences may make it less valid to compare or combine the results of these studies.

Employment characteristics were under-reported in all three studies, as can be seen in Table 120.

Table 118. Internal validity of studies of splinting after carpal tunnel release

Study	Number of patients	Percent bilateral patients	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
Finsen, 1999 319	74	10.8%	Single	Not reported	RCT	No	0	Yes	92.7
Bury, 1995 364	47	7.5%	Single	Not reported	RCT	No	7	No	NR
Cook, 1995 432	50	0%	Single	Not reported	RCT	No	0	Yes	NA

Table 119. Generalizability of studies of splinting after carpal tunnel release

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Finsen, 1999 319	74	54.7 (21-86)	81.1	NR	0	0	0	0	0	0	0	No	No
Bury, 1995 364	47	41.4 (19-79)	83.0	13 (5-36)	NR	NR	NR	7	NR	NR	NR	No	Yes
Cook, 1995 432	50	NR	NR	NR	0	NR	0	NR	0	0	0	Yes	Yes

Table 120. Patient employment characteristics in studies of splinting after carpal tunnel release

Study	Number of patients	% Patients employed	% Patients receiving workers' compensation	% Patients retired	% Patients Homemakers	Reported occupations
Finsen, 1999 319	74	63.5	Not reported	Not reported	Not reported	Not reported
Bury, 1995 364	47	Not reported	Not reported	Not reported	Not reported	Not reported
Cook, 1995 432	50	Not reported	16.0	Not reported	Not reported	Not reported

Table 121. Effect of splinting after surgery on global outcome

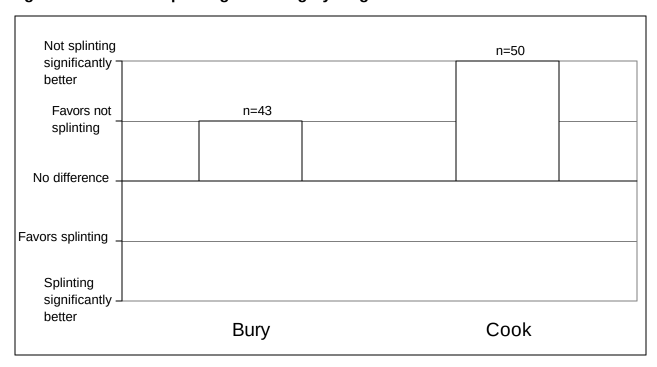
Study	Number of Patients	Global Outcome	Statistical Significance of Difference Between Groups
Bury et al., 1995 364		Global score (Scale not reported)	Not reported
	No splint 17	8.0	
	2 week splint 26	8.1	
		Number of patients symptom free	Not significantly different by chi square test conducted by ECRI, p = 0.85.
	No splint 17	9	
	2 week splint 26	13	
		Categorical rating	Not significantly different by chi square test conducted by ECRI, p = 0.68.
	No splint 17	Cured: 8 Improved: 9 Unchanged: 0 Worse: 0	Not significantly different when data is collapsed into a dichotomous outcome (number cured or improved) by chi square test conducted by ECRI, p = 0.15
	2 week splint 26	Cured: 12 Improved: 11 Unchanged: 1 Worse: 2	conducted by Estat, p. 10.10
Cook et al., 1995 432		14 Days:	
	No splint 25	Excellent 9 Good 9 Fair 7	Significantly different by chi square test
	2 week splint 25	Excellent 1 Good 14 Fair 10	conducted by ECRI, p = 0.018.
		1 Month:	
	No splint 25	Excellent 12 Good 10 Fair 3	Significantly different by Chi square test conducted by ECRI, p = 0.007.
	2 week splint 25	Excellent 2 Good 18 Fair 5	

Table 122. Summary of effect of splinting after surgery on global outcome

Study	Which Procedure led to Superior Global Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Bury, 1995 364	No Splint	No	Number symptom free 28% Categorical rating ^b 29%	Number symptom-free 0.06 (-0.61 – 0.72) Categorical rating ^b 0.89 (-0.78-2.56)
Cook, 1995 432	No Splint	Yes	Not calculable	14 Days 0.38 (-0.18-0.94) 1 Month 0.86 (0.28-1.44)

a: Calculated by ECRI

Figure 41. Effect of splinting after surgery on global outcome



b: Calculated by ECRI by collapsing the categorical rating into a dichotomous one: number cured or improved.

Return to work

All three trials reported on return to work. These results are presented in Table 123. As can be seen in Table 124 and Figure 42, two studies show a trend toward favoring no splint, with the difference becoming statistically significant in the study by Cook. In contrast, the study by Finsen shows no difference between groups.

Table 123. Effect of splinting after surgery on return to work

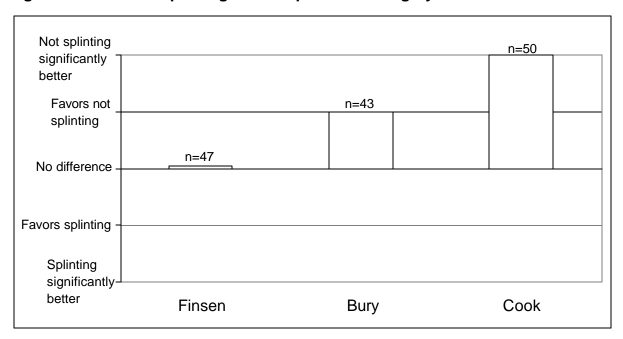
Study	Number of Patients	Return to work	Statistical Significance of Difference Between Groups
Finsen, 1999 319	No colint 20	Median time to return to work	Not reported
	No splint 28 4 week splint 19	6 Weeks (95% CI 5-6 Weeks) 6 Weeks (95% CI 4-7 Weeks)	
Bury, 1995 ³⁶⁴	No splint 17 2 week splint	Numbera of patients who had not returned to work at last followup (Mean 5.7 Months) 2	Not significantly different by chi-square test conducted by ECRI, p = 0.23
0 1 1005 100	26	7	
Cook, 1995 ⁴³²	No splint 25	Time to return to work Light duty: 15 Days Full duty: 17 Days	Significantly different by t-test (Light duty p = 0.01; Full duty p = 0.005)
	2 week splint 25	Light duty: 24 Days Full duty: 27 Days	

a: Calculated by ECRI from a published percentage

Table 124. Summary of effect of splinting after surgery on return to work

Which Procedure led to Superior Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
No difference	No	Not calculable	Not calculable
No Splint	No	24%	0.55 (-0.39 – 1.49)
No Splint	Yes	Not calculable	Light duty: 0.75 (0.17 – 1.32) Full duty: 0.82 (0.24-1.40)
	Procedure led to Superior Outcome? No difference	Procedure led to Stastically Significant? No difference No No Splint No	Procedure led to Stastically Significant? No difference Outcome? No Splint Difference Stastically Significant? No Splint No 24% (Minimum percent difference detectable) No Not calculable

Figure 42. Effect of splinting after carpal tunnel surgery on return to work



Return to Activities of Daily Living

One study of 50 patients reported on time to return to activities of daily living. The results are presented in Table 125. These results show a statistically significant advantage to not splinting. The effect size is significantly different from zero (d = 1.06, 95% C.I. 0.47 - 1.65).

Table 125. Effect of splinting after surgery on time to return to activities of daily living

Study	Number of Patients	Return to Activities of Daily Living	Statistical Significance of Difference Between Groups
Cook, 1995 432		Time to return to activities of daily living	Significantly different by Hest, p = 0.0004.
	No splint 25	6 Days	
	2 week splint 25	12 Days	

<u>Pain</u>

Two studies reported on pain. The results are presented in Table 126. Finsen et al. found no statistically significant differences between groups. Cook et al. found statistically significant differences between groups at 2 weeks and 4 weeks. These differences were stated to be no longer significant at 3 and 6 months, but no data were reported. In this study, it is unclear whether the pain described after treatment is pain from carpal tunnel syndrome, pain resulting from surgery, or both. As can be seen in Table 127 and Figure 43, the results of the two studies show opposite trends, and as noted above, it is unclear whether the patients in these two studies are comparable.

Table 126. Effect of splinting after surgery on pain

Study	Number of Patients	Pain	Statistical Significance of Difference Between Groups
Finsen, et		Median VAS (0-100)	Not significantly different (p >0.05; test not
al., 1999 319	No splint 45		reported)
	Preop	56 (Range 46-65)	
	2 Weeks	6 (Range 4-17)	
	6 Months	3 (Range 2-8)	
	4 week splint 37		
	Preop	51 (Range 38-57)	
	2 Weeks	5 (Range 2-11)	
	6 Months	2 (Range 0-4)	
Cook et al.,		Verbal Scale (1-10)	Significantly different at both time points
1995 ⁴³²	No splint 25		(p = 0.001 and 0.01 respectively by t-test)
	14 Days	0.9	
	1 Month	0.5	
	2 week splint 25		
	14 Days	2.4	
	1 Month	1.5	

Table 127. Summary of effect of splinting after surgery on pain

Study	Which Procedure led to Superior Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Finsen, et al., 1999 319	Splinting	No	Not calculable	Not calculable
Cook et al., 1995 ⁴³²	No Splint	Yes	Not calculable	14 Days: 0.98 (0.39 – 1.56) 1 Month: 0.75 (0.17 – 1.32)

a: Calculated by ECRI

Table 128. Reported harms in studies of splinting after carpal tunnel surgery

Study	Patients per group	Complication	Number reporting
Finsen, 1999 319	No splint 45	Superficial Hematoma	1
		Wound discharge	1
	2 Week splint 36	Superficial Hematoma	0
		Wound discharge	0
Bury, 1995 ³⁶⁴	No splint 17	Persistent symptoms requiring	1
		reoperation	
	2 week splint 26	Persistent symptoms requiring	0
		reoperation	
Cook, 1995 432	No splint 25	Reported that there were no wound	0
		complications or bowstringing tendons	
	2 week splint 25		

Conclusions

The three studies examining whether there was an advantage to splinting after carpal tunnel surgery have yielded fairly consistent results within each study. Cook, et al found a statistically significant advantage to not splinting for reduced pain, faster return to work and daily activities, and superior global outcome. Bury also found that not splinting led to better global outcome and faster return to work, but neither of these effects was statistically significant. This study lacked the statistical power to detect small (<20%) differences between groups. In contrast, Finsen et al. found a small and statistically nonsignificant trend advantage for the effect of splinting on pain, while times to return to work were the same for both groups. The reasons for the differences between studies is not readily apparent from an examination of the study or patient characteristics. There may be conditions under which splints offer an advantage and conditions under which they do not. Further studies are necessary before a conclusion may be reached.

What are the relative benefits and harms of vitamin B therapy for persons with carpal tunnel syndrome?

One trial of 17 patients examining the effect of vitamin B_6 therapy on carpal tunnel syndrome met exclusion criteria. 459

Internal Validity

This was a small (n = 15) randomized controlled trial. There was 13% attrition, and compliance was not reported. Study characteristics affecting internal validity are listed in Table 129.

Generalizability

This study did not report patient characteristics except that patients with mild disease were excluded, so no discussion of its generalizability is possible.

Table 129. Internal validity of studies of vitamin B therapy for carpal tunnel syndrome

Study	Number of patients	Number of centers	Funded by a for-profit stakeholder?	Study design	Blinding	Total Patient Attrition (all patient groups)	Intent to treat analysis?	% Compliance
Stransky, 1989 ⁴⁵⁹	15	Single	Not reported	RCT	Double	2	No	NR

Table 130. Generalizability of studies of vitamin B therapy for carpal tunnel syndrome

Study	Number of patients	Mean age and range	% Female	Mean Duration (Months) of condition and range	% Patients with diabetes	% Patients with arthritis	% Patients with previous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy		% Patients on kidney dialysis	Excluded severe disease?	Excluded mild disease?
Stransky, 1989 459	15	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	No	Yes

Results

This trial reported a single patient-oriented outcome (global outcome expressed as number of patients improved after treatment). A summary of the effect of vitamin B_6 therapy in this study is shown in Table 131. There was no statistically significant difference in percent of patients improved between-treatment groups. This study had few patients and very low power. Only large (46-48%) differences between groups were would have been statistically significant.

Table 131. Global outcome in patients treated with vitamin B therapy

Study	N (units)	Global outcome - number (%) patients improved	Statistical significance of difference between groups
Stransky et al. 1989	Vitamin B ₆ 6	3 (50)	Vitamin B ₆ was not significantly
459			different from placebo or control by
	Placebo 5	4 (80)	chi-square test conducted by ECRI
			(p = 0.30 and 0.42, respectively)
	Untreated Control 4	3 (75)	

Table 132. Summary of effect of vitamin B therapy on symptoms of carpal tunnel syndrome

Study	Which Treatment led to Superior Global Outcome?	Was the Difference Stastically Significant?	Power (Minimum percent difference detectable) ^a	Effect Size (95% Confidence Interval) ^a
Stransky et al. 1989 ⁴⁵⁹	Placebo	No	Vitamine vs. Placebo 46%	Vitamine vs. Placebo -0.55 (-1.86 – 0.75)
			Vitamine vs. No treatment	Vitamine vs. No treatment
			48%	-0.42 (-1.76 – 0.91)

a: Calculated by ECRI

Conclusions

Although the low power of the study prevents any solid conclusion from being drawn, the trend toward a greater percentage of improved patients in the placebo group does not support the therapeutic effectiveness of Vitamin B_6 .

Question #4: Is there a relationship between specific clinical findings and specific treatment outcomes among patients with carpal tunnel syndrome?

In addressing this question, we consider whether published literature suggests that there are clinical findings that predict positive or negative outcomes after treatment for carpal tunnel syndrome. The studies we considered all attempted to identify predictors by using regression techniques or by comparing outcomes in different groups of patients with different pre-treatment clinical findings.

Excluded studies

As discussed in the Methods section, we retrieved articles identified by our literature searches according to certain *a priori* criteria. However, not all of the retrieved studies met our more specific inclusion criteria for this question. These latter studies, and the reason we did not consider them for this question are shown in Table 133.

Table 133. Excluded studies

Author	Reason for exclusion
Walker (2000)	Stratified study that did not examine any correlations that were
34	also examined by at least two other studies
Hasegawa	Stratified study that did not examine any correlations that were
(1999) ³⁴	also examined by at least two other studies
Olney (1999)	Stratified study that did not examine any correlations that were
323	also examined by at least two other studies
Rosen (1997)	Stratified study that did not examine any correlations that were
343	also examined by at least two other studies
LoMonaco	Stratified study that did not examine any correlations that were
(1996) ³⁵⁸	also examined by at least two other studies
Padua (1996)	Stratified study that did not examine any correlations that were
358	also examined by at least two other studies
Wintman	Stratified study with no clinical finding/outcome comparisons
(1996) ³⁶²	reported by at least three studies
Chang and	Stratified study that did not examine any correlations that were
Dellon (1993)	also examined by at least two other studies
389	

Evidence base

After these exclusions, there remained 12 studies with a total of 1723 patients.

Study quality

The evaluation of the quality of literature for this question differs from quality evaluations of studies of treatments. This is because, for the present question, the RCT is not necessarily the "gold standard". Case series data, if appropriately analyzed, can also yield valid information. Consequently, the method of data analysis plays a prominent role when considering the quality of the studies relevant to this question.

Table 134. Study quality

Author/year	Prospective?	Methods used to identify predictor variables
Finsen and Russwurm (2001) ²²⁴	Yes	Stratification
Shin (2000) 460	No	Multiple logistic regression
Straub (1999) 305	Yes	Stratification
Atroshi (1998) 461	Yes	Multiple linear regression
Choi and Ahn (1998) 329	No	Stratification
Katz (1998) 462	Yes	Multiple logistic regression
Higgs (1997) 341	No	Stratification
Glowacki (1996) 352	No	Stratification
Jacobsen and Rahme (1996) 353	Yes	Multiple regression ^a
Al-Qattan (1994) 375	No	Stratification
Nathan (1993) 395	Partly ^b	Multiple regression
Yu (1992) ⁴⁰³	No	Stratification

alndependent analysis of individual patient data conducted by ECRI

bPatients entering the study after a certain date were studied prospectively; patients who had treatment prior to that date were studies retrospectively.

Results

Table 135 shows the relationship of specific clinical findings to treatment outcomes in those studies that used regression to identify predictor variables. In the table, clinical variables are indicated by boldface type. There are five such studies with a total of 932 patients. Also presented in this table are non-clinical variables (e.g. age, gender) to show all of the variables used in each multiple regression.

No study that employed regression analysis reported statistically significant correlations between two-point discrimination or grip strength and any outcomes. However, three out of four studies that examined the "predictability" of electrodiagnostic tests reported statistically significant correlations between electrodiagnostic test results and various outcomes. Two of the studies that found a statistically significant relationship were prospective.

The outcomes predicted by electrodiagnostic test results in the three "significant" studies were odds of obtaining disability payment, patient satisfaction with surgery, and number of sick leave days. Odds of obtaining disability payment were higher in patients diagnosed with CTS (mild, moderate, or severe) compared to those with normal electrodiagnostic findings. Another study found patient satisfaction with surgery was lower among patients with a better electrodiagnostic test (distal motor latency) before surgery. Analysis of individual patient data from a third study revealed that number of sick leave days was higher among patients with a pre-surgical electrodiagnostic test indicating slight or intermediate CTS as opposed to pronounced CTS. In the fourth

Table 135. Relationship between specific clinical findings and treatment outcomes among patients with Carpal Tunnel Syndrome (Multiple regression analysis)

Author	N	Treatment	Outcomes	٧						st two s	studies me?)	i	Unique study variables
				Age	Gender	Treatment	Hand dominance	Insurance type	Employment status	Two-point discrimination ^d	Electrodiagnostic test	Grip strength	
Shin (2000) 460	210	Conservative treatments Surgery ^a	Odds of obtaining employment disability	NS	NS	NS	_	_	_	_	Sig	_	Mechanism of injury (NS)
Atroshi (1998) ⁴⁶¹	140	Surgery ^b	Global outcome (patient dissatisfaction)	Sig	NS	_	NS	_	NS	NS	Sig	NS	Vibration exposure (sig), ADL score (NS), thenar atrophy (NS), pinch strength (NS), tinel sign (NS), phalen sign (NS)
Katz (1998) 462	315	Surgery and conservative treatments (not described)	Work absence (18 months after treatment)	NS	NS	NS	_	NSe	NS	-	-	NS	Occupation (NS), baseline function (sig), function at 6 months (sig), hired attorney (sig), work absence at enrollment (NS), work absence at 6 months (sig), mental health status (NS), physical and clerical self-reported exposure scales
Jacobsen and Rahme (1996) 353	29 (32 hands)	Surgery ^c	Number of sick days after surgery	NS	NS	NS	NS	_	_	NS	Sig	-	None
Nathan (1993) ³⁹⁵	238	Surgerya	Return to work	NS	NS	_	NS	Sig	NS	_	NS	-	Laterality (NS), year of study (NS), referral source (NS), incision length (NS), occupational hand use (NS), diabetes (NS),

Author	N	Treatment	Outcomes	Variables examined by at least two studies (significant correlation with outcome?)							Unique study variables		
				Age	Gender	Treatment	Hand dominance	Insurance type	Employment status	Two-point discrimination ^d	Electrodiagnostic test	Grip strength	
													rheumatoid arthritis (NS), number and density of hand therapy sessions/ week (NS)

aOpen release
bUnilateral endoscopic release
cOpen and endoscopic release
dVariables in boldface represent clinical findings
dIn a related publication, surgical patients alone were analyzed and insurance type significantly correlated with work absence 6 months post-surgery.302
NS – Not significant

Table 136. Stratified studies (global outcome)

Study	N	Treatment	Stratification variable Electrodiagnostic nerve deficit			
Finsen and Russworm (2001) ²²⁴	79	Surgery (open release)	VAS for pain and discomfort	NS		
Straub (1999) 305	100	Surgery (endoscopic release)	Satisfactory/unsatisfactory result	NS (but trend toward more success in abnormal sensory/ normal motor nerve conduction group)		
Choi and Ahn (1998)	154	Surgery (open release)	Patient satisfaction (poor, fair, good, or excellent)	NS		
Higgs (1997)	93	Surgery (open release)	Improved/not improved	Sig (normal/near normal)		
Glowacki (1996) 352	167	Surgery (open release)	Symptoms resolved, improved, or same or worse	NS		
Al-Qattan (1994) 375	112	Surgery (open release)	Satisfactory/poor outcome	NS		
Yu (1992) 403	53	Surgery (open release)	Good/fair/poor result	NS		

NS – Not signficant

Conclusions

Studies that searched for relationships between clinical findings and treatment outcomes did so by using multiple regression analysis or stratified patient groups. Among studies that used regression analysis, the only clinical finding variable shown by more than one study to significantly predict treatment outcomes was electrodiagnostic testing. This finding was statistically significant in three of the four studies that examined it. The outcomes predicted by these three studies were patient satisfaction with surgery, odds of obtaining disability payment, and number of sick days after surgery. Odds of obtaining disability payment were higher in patients diagnosed with CTS (mild, moderate, or severe) compared to those with normal electrodiagnostic findings. Another study found patient satisfaction with surgery was lower among patients with a better electrodiagnostic test results (distal motor latency) before surgery. Analysis of individual patient data from a third study revealed that number of sick leave days was higher among patients with a pre-surgical electrodiagnostic test indicating slight or intermediate CTS as opposed to pronounced CTS. The fourth study of electrodiagnostic tests found no statistically significant relationship between electrodiagnostic test results and return to work. This apparent lack of consistency of results could indicate that, although the relationship between electrodiagnostic tests and treatment outcomes is statistically significant, it may not be substantial. The possibility that this relationship is small is supported by the results of stratified studies that examined the relationship between electrodiagnostic test results and global outcomes. Six of seven studies did not find a statistically significant relationship.

Question #5: Is there a relationship between duration of symptoms and specific treatment outcomes among patients with carpal tunnel syndrome?

In addressing this question, we consider whether published literature suggests that duration of symptoms predicts positive or negative outcomes after treatment for carpal tunnel syndrome. The studies we considered all attempted to identify predictors by using regression techniques or by comparing outcomes in different groups of patients with different duration of symptoms.

Excluded studies

As discussed in the Methods section, we retrieved articles identified by our literature searches according to certain *a priori* criteria. However, not all of the retrieved studies met our more specific inclusion criteria for this question. These latter studies, and the reason we did not consider them for this question are shown in Table 137.

Table 137. Excluded studies

Author Reason for exclusion								
Wintman	Stratified study with no duration of symptoms/outcome							
(1996) ³⁶²	comparisons reported by at least three studies							

Evidence base

After this exclusion, there remained six studies with 984 patients.

Study quality

The criteria used to evaluate study quality were identical to those described for Question 4. One prospective study and one retrospective study conducted a multiple regression analysis, while four studies performed stratifications

(Table 138). Only one of the four stratified studies was prospective in design.

Table 138. Study quality

Author/year	Prospective?	Methods used to identify predictor variables
Straub (1999) 305	Yes	Stratification
Atroshi (1998) 461	Yes	Multiple linear regression
Choi and Ahn (1998) 329	No	Stratification
DeStefano (1997) 463	No	Multivariable proportional hazards regression
Al-Qattan (1994) 375	No	Stratification
Yu (1992) ⁴⁰³	No	Stratification

Table 139. Relationship between duration of symptoms and treatment outcomes among patients with Carpal Tunnel Syndrome.

Author	N	Treatment	Outcomes	Duration of symptoms – significance (duration associated with better outcome)	Other variables examined
Atroshi (1998) ⁴⁶¹	140	Surgery (unilateral endoscopic release)	Global outcome (patient dissatisfaction)	NS	Age (sig), sex (NS), hand dominance (NS), unemployment (NS), vibration exposure (sig), ADL score (NS), DML (sig), surgeon (NS), subjective weakness (NS), type of work (NS), type of symptoms (NS), Tinel sign (NS), Phalen's test results (NS), thenar atrophy (NS), two-point discrimination (NS), grip strength (NS), pinch strength (NS)
DeStefano (1997) ⁴⁶³	425	Non-surgical (oral meds, oral steroids, steroid injections, splints) Surgical (carpal tunnel release)	Global outcome (symptom resolution)	NS (non-surgical patients) Sig (surgical patients, <3 years)	Age (NS), sex (NS), carpal tunnel syndrome category (NS), hand involved (NS), arthritis (NS), pregnancy (NS), injury (NS), diabetes or hypothyroidism (sig for surgical patients)

Table 140. Stratified studies (global outcome)

Study	N	Treatment	Global outcome measure	Duration of symptoms – significance (duration associated with better outcome)
Straub (1999) 305	100	Surgery (endoscopic release)	Satisfactory/unsatisfactory result	NS
Choi and Ahn (1998) 329	154	Surgery (open release)	Patient satisfaction (poor, fair, good, or excellent)	Sig (shorter duration, <3 months)
Al-Qattan (1994) 375	112	Surgery (open release)	Satisfactory/poor outcome	NS
Yu (1992) 403	53	Surgery (open release)	Good/fair/poor result	NS, but trend toward more success in ≥6 month group

NS – Not signficant

Question #6: Is there a relationship between factors such as patients' age, gender, socioeconomic status and/or racial or ethnic grouping and specific treatment outcomes among patients with carpal tunnel syndrome?

In addressing this question, we consider whether published literature suggests that there are demographic variables that predict positive or negative outcomes after treatment for carpal tunnel syndrome. The studies we considered all attempted to identify predictors by using regression techniques or by comparing outcomes in different groups of patients with different pre-treatment demographic characteristics.

Excluded studies

As discussed in the Methods section, we retrieved articles identified by our literature searches according to certain *a priori* criteria. However, not all of the retrieved studies met our more specific inclusion criteria for this question. These latter studies, and the reason we did not consider them for this question are shown in Table 141.

Table 141. Excluded studies

Author	Reason for exclusion
Walker (2000) 34	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Braun (1999) 316	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Hasegawa (1999) ³⁴	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Higgs (1997) 341	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Rosen (1997) 343	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Padua (1996) 358	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Wintman (1996) 362	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Nancollas (1995) 464	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Chang and Dellon (1993) 389	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies
Feinstein (1993) 390	Data presentation did not allow determination of correlation
Hagberg (1991) 308	Stratified study with no demographic variables/outcome
	comparisons reported by at least three studies

Table 142. Study quality

Author/year	Prospective?	Methods used to identify predictor variables
Shin (2000) 460	No	Multiple logistic
01 (1000) 000		regression
Olney (1999) 323	No	Stratification
Straub (1999) 305	Yes	Stratification
Atroshi (1998) 461	Yes	Multiple linear
		regression
Davies (1998) 330	No	Stratification
Katz (1998) 462	Yes	Multiple logistic
		regression
DeStefano	No	Multivariable
(1997) ⁴⁶³		proportional hazards
		regression
Elmaraghy and	Yes	Stratification
Hurst (1996) 349		
Glowacki (1996) 352	No	Stratification
Jacobsen and Rahme (1996) 353	Yes	Multiple regression
Lee and Jackson (1996) 355	No	Stratification
Nagle (1996) 357	Yes	Stratification
Strickland (1996) 361	No	Stratification
Wintman (1996) 362	Yes	Stratification
Hallock and Lutz (1995) 368	Yes	Stratification
Mirza (1995) 371	Unknown	Stratification
Al-Qattan (1994)	No	Stratification
Roth (1994) 383	Yes	Stratification
Nathan (1993) 395	Partly ^a	Multiple regression
Palmer (1993) 397	Yes	Stratification
Agee (1992) 46	Yes	Stratification
Yu (1992) ⁴⁰³	No	Stratification

WC – Workers' compensation

bPatients entering the study after a certain date were studied prospectively; patients who had treatment prior to that date were studies retrospectively

Results

Table 143 shows the relationship of specific demographic variables to treatment outcomes in those studies that used regression to identify predictor variables (demographic variables are shown in bold type). There are six such studies with a total of 1357 patients. Also presented in this table are non-demographic variables (e.g. grip strength) to show all of the variables used in each multiple regression.

Table 143. Relationship between demographic factors and treatment outcomes among patients with Carpal Tunnel Syndrome (multiple regression analysis)

Author	N	Treatment	Outcomes	V	Variables examined by at least two studies (significant correlation with outcome?)								Unique study variables
				Age	Gender	Treatment	Hand dominance	Insurance type	Employment status	Two-point discrimination	Electrodiagnostic test	Grip strength	
Shin (2000) ⁴⁶⁰	210	Conservative treatments Surgery ^a	Odds of obtaining employment disability	NS	NS	NS	_	_	_	-	Sig	_	Mechanism of injury (NS)
Atroshi (1998) ⁴⁶¹	140	Surgeryb	Global outcome (patient dissatisfaction)	Sig	NS	_	NS	_	NS	NS	Sig	NS	Vibration exposure (sig), ADL score (NS), thenar atrophy (NS), pinch strength (NS), tinel sign (NS), phalen sign (NS)
Katz (1998) ⁴⁶² ³⁰²	315	Surgery and conservative treatments (not described)	Work absence (18 months after treatment)	NS	NS	NS	1	NS (all patients) e Sig (surgery patients)	NS	NS	-	NS	Occupation (NS), baseline function (sig), function at 6 months (sig), hired attorney (sig), work absence at enrollment (NS), work absence at 6 months (sig), mental health status (NS), physical and clerical self-reported exposure scales
DeStefan o (1997) 463	425	Conservative treatments Surgery (carpal tunnel release)	Global outcome (symptom resolution)	NS	NS	Si g	NS	_	_	Sig (sur gica I pati ents only)	Ι	1	_

Author	N	Treatment	Outcomes	V	Variables examined by at least two studies (significant correlation with outcome?)							3	Unique study variables
				Age ^d	Gender	Treatment	Hand dominance	Insurance type	Employment status	Two-point discrimination	Electrodiagnostic test	Grip strength	
Jacobsene and Rahme (1996) 353	29 (32 hands)	Surgery ^c	Number of sick days after surgery	NS	NS	NS	NS	-	_	NS	Sig	_	None
Nathan (1993) ³⁹⁵	238	Surgerya	Return to work	NS	NS	_	NS	Sig	NS	_	NS	_	Laterality (NS), year of study (NS), referral source (NS), incision length (NS), occupational hand use (NS), diabetes (NS), rheumatoid arthritis (NS), number and density of hand therapy sessions/ week (NS)

aOpen release
bUnilateral endoscopic release
cOpen and endoscopic release
dVariables in boldface represent demographic characteristics
eln a related publication, surgical patients alone were analyzed and insurance type significantly correlated with work absence 6 months post-surgery.³⁰²
eMultiple regression performed independently by ECRI from individual patient data presented in this study
NS – Not significant

Table 144. Stratified studies (global outcome)

Study	N	Treatment	Global outcome	Stratification	variable
			measure	Workers'	Job
				compensation (WC) status	category
Straub (1999) 305	100	Surgery (endoscopic release)	Satisfactory/unsatisfactory result	NS (but trend toward more success in non- WC group)	NS
Davies (1998) 330	239	Surgery (endoscopic release)	Patient satisfaction/dissatisfaction	Sig (non-WC)	_
Glowacki (1996) 352	167	Surgery (open release)	Symptoms resolved, improved, or same or worse	Sig (non-WC)	_
Al-Qattan (1994) 375	112	Surgery (open release)	Satisfactory/poor outcome	Sig (non-WC)	Sig (not physically strenuous)
Yu (1992)	53	Surgery (open release)	Good/fair/poor result	_	Sig (not physically strenuous)

NS – Not significant

Table 145. Stratified studies (return to work)

Study	N	Treatment	Stratification variable
			Workers'
			compensation (WC) status
Olney (1999)	211	Surgery (open or endoscopic release)	Sig (non-WC and non- contested WC)
Davies (1998)	239	Surgery (endoscopic release)	Sig (non-WC)
Elmaraghy and Hurst (1996) 349	75	Surgery (endoscopic release)	Sig (non-WC)
Lee and Jackson (1996) 355	237	Surgery (limited incision release using carposcope)	Sig (non-WC)
Nagle (1996) 357	291	Surgery (endoscopic release)	Sig (non-WC)
Strickland (1996) ³⁶¹	62	Surgery (hypothenar fat pad flap for patients who received unsuccessful open release)	NS, except for manual labor subgroup (non-WC)
Hallock and Lutz (1995) 368	96	Surgery (open or endoscopic release)	Sig (non-WC)
Mirza (1995) 371	236	Surgery (endoscopic release)	Sig (non-WC)
Roth (1994) 383	95	Surgery (endoscopic release)	Sig (non-WC)
Palmer (1993) ³⁹⁷	163	Surgery (open or endoscopic release)	Sig (non-WC)

Study	N	Treatment	Stratification variable
			Workers' compensation (WC) status
Agee (1992)	122	Surgery (open or endoscopic release)	Sig (non-WC)

NS – Not significant

Conclusions

The available evidence suggests that patients who are not receiving workers' compensation tend to return to work faster than those receiving such compensation. This is suggested by one of two "multiple regression" studies of this relationship and by a combination of 10 prospective and retrospective stratified studies. Some evidence also suggests that patients who are not receiving workers' compensation have better global outcomes, but this evidence is derived exclusively from retrospective studies. Therefore, these latter findings require confirmation. In any event, one cannot ascribe causal relationships to these correlations.

Available evidence suggests that there is no strong relationship between gender, employment status, or hand dominance and return to work or global outcomes. There is insufficient evidence to arrive at a firm evidence-based conclusion on the relationship between type of work, diabetes, or age and patient outcomes.

We define an instrument that can accurately assess functional limitations in an individual with carpal tunnel syndrome as one that has been shown to have: test-retest reliability, internal reliability, concurrent validity, predictive validity, and responds to treatment.

Table 146. Potential biases in assessment instruments^a

Bias	Definition
Yea-saying	The tendency to always agree with yes-no questions.
End aversion	The tendency to use middle values rather than the end points of analog scales
Question framing	The tendency for the wording of a question to affect the response.
Motivation to seem better	Patients want to subconsciously please their health-care providers by responding to
	treatment and are embarrassed to complain about problems.
Motivation to seem worse	Can occur if patients will lose services or benefits if they improve.
Response shifts	The tendency of patients to modify their internal standards of evaluation so that their
	current level of functioning is seen as normal.
Memory failure	Difficulty in remembering past function may influence assessment of current function.
Leading the patient	The tendency of the questionnaire itself to change the way the patient assesses
	functioning.

a Adapted from Gotay 1996474

Evidence base

Eight studies met the inclusion criteria (see the section Inclusion Criteria). They are listed in Table 147. The functional assessment instruments evaluated by the studies that met the inclusion criteria are listed in Table 148.

Table 147. Trials of functional assessment instruments that met the inclusion criteria

Study	Instruments evaluated ^a	N subjects	Outcome measurements
Vaile 1999 475	NHP, SF-36, mSHAQ, V-VAS	27	Response to treatment
Alderson 1999 315	AMHFQ	26	Validity Test-retest reliability
Atroshi 1998 326	SF-36 and CTS-I	102	Test-test comparison Test-retest reliability Response to treatment
Pransky 1997 476	UEF	165	Validity Test-test comparison
Atroshi 1997 477	SF-36 and CTS-I	277	Validity
Katz 1994 377	Global score	104	Validity
Katz 1994 303	CTS-I and K-ADL	74	Response to treatment
Levine 1993 ³⁹³	CTS-I	67	Validity Test-retest reliability Response to treatment

a The full names of the instruments and descriptions of the instruments are given in

Table 148. Instruments evaluated to measure functional limitations associated with carpal tunnel syndrome

Instrument	Abbreviation	First described by	Scoring system	Subjects covered	Extent of use ^a
Alderson-McGall Hand Function Questionnaire	AMHFQ	Alderson and McGall 1999 ³¹⁸	Functional difficulty categories	Common tasks performed with the hands	Not widely used
Calculated Global Score	Global Score	Katz 1994 ³⁷⁷	VAS	Grip strength, numbness, pain, parethesia	Not widely used
Carpal Tunnel Syndrome Instrument	CTS-I	Levine 1993 ³⁹³	Functional difficulty categories/ symptom severity categories	Eight ADL, and severity of symptoms	Widely used
Katz Activities of Daily Living	K-ADL	Katz 1994 ³⁰³	Functional difficulty categories	Ten ADL	Not widely used
Medical Outcomes Study 36-Item Short-Form Health Survey	SF-36	Ware 1992 ⁴⁷⁸	Categories	Impact of health on physical activities, social activities, activities of daily living, pain, psychological distress, emotional health, and energy	Extensively used
Modified Stanford Health Assessment Questionnaire	mSHAQ	479	Categories	ADL	Widely used
Nottingham Health Profile	NHP	Hunt 1985 ⁴⁸⁰	Categories	Pain, energy, emotional reactions, sleep problems, social isolation, physical mobility, employment, hobbies, sex life, personal relationships, and holiday	Widely used
Upper Extremity Function Scale	UEF	Pransky 1997 ⁴⁷⁶	Functional difficulty categories	Eight ADL	Not widely used
Vaile Visual Analog Scales	V-VAS	Vaile 1999 ⁴⁷⁵	VAS	Impact of CTS on well being, discomfort, activities	Not widely used

^aExtent of use was determined by searching Medline for manuscripts that used the assessment instrument. Not widely used = 3 or fewer studies. Widely used= four to ten studies. Extensively used= more than ten studies.

Table 149. Details of study design

Study	Number of patients	Number of centers	Funded by a for-profit agency?	Study design	Prospective	Blinding	% Attrition	Intent to treat analysis	% Compliance
Vaile 1999 475	27	2	NR	Cohort	Yes	No	0	Yes	NA
Alderson 1999 315	26	1	NR	Cohort	Yes	Rater	34	No	NA
Atroshi 1998 326	102	1	No	Cohort	Yes	Rater	0	Yes	NA
Pransky 1997 476	165	1	No	Cohort	Yes	No	44.8	No	NA
Atroshi 1997 477	277	3	No	Cohort	Yes	No	23.4	No	NA
Katz 1994 377	104	4	No	Cohort	Yes	Rater	0	Yes	NA
Katz 1994 303	74	4	NR	Cohort	Yes	Rater	NR	No	NA
Levine 1993 393	67	2	No	Cohort	Yes	Rater	0	No	NA

Generalizability

It is important for studies that evaluate assessment instruments to enroll patients who are representative of the population of interest. Information about patients enrolled in the studies addressing this question are shown in Table 150. All eight studies recruited populations that appear to be "typical" of patients presenting with carpal tunnel syndrome as has been established by epidemiology studies (See the Introduction). The patient groups are predominantly female and middle aged. Few of the studies reported on the presence of co-morbid conditions that may have contributed to functional limitations. The occupations and employment status of the patients are shown in Table 151. The two studies by Katz recuited patients from the same large randomized controlled trial, a trial that was comparing different methods of surgically treating carpal tunnel syndrome.

Table 150. Study generalizability: patient characteristics

Study	Number of patients	Mean age and range	% female	Duration of conditon mean and range months	% Patients with diabetes	% Patients with arthritis	% Patients with prevous relevant injuries	% Patients with other relevant nerve impingement conditons	% Patients with peripheral neruopathy	% Patients pregnant	% Patients on kidney dialysis	Did the study exclude patients with severe disease?	Did the study exclude patients with mild disease?
Vaile 1999 475	27	57 (29-84)	81.4	NR	NR	55.5	NR	NR	NR	NR	NR	No	No
Alderson 1999 315	26	44.4 (22-79)	70.5	(3-48)	NR	0	NR	NR	NR	NR	NR	No	No
Atroshi 1998 326	102	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
Pransky 1997 476	165	46 (19-65)	67	NR	NR	NR	NR	NR	NR	NR	NR	No	No
Atroshi 1997 477	277	46.6 (13-91)	77.8	NR	NR	NR	NR	NR	NR	NR	NR	No	No
Katz 1994 377	104	55 (25-87)	70	NR	NR	NR	NR	NR	NR	NR	NR	No	No
Katz 1994 ³⁰³	74	55 (25-87)	70	NR	NR	0	NR	0	0	0	NR	No	No
Levine 1993 ³⁹³	67	57 (19-88)	75	18 (3-58)	NR	NR	NR	NR	NR	NR	NR	NR	No

Table 151. Generalizability: employment status and occupations

Study	Number of patients	% Patients employed	% Patients on Workers	% Patients retired	% Patients homemakers	Reported Occupations
Vaile 1999 475	27	NR	0	NR	NR	NR
Alderson 1999 315	26	NR	35	NR	5.6%	Business-17.6% Sciences-5.9% Health-11.8% Education-5.9% Recreation-5.9% Sales-11.8% Trades and Transport-5.9% Industry-5.9% Manufacturing-23.5%
Atroshi 1998 326	102	NR	NR	NR	NR	NR
Pransky 1997 ⁴⁷⁶	165	89	10	NR	NR	NR
Atroshi 1997 477	277	NR	28.8	NR	NR	NR
Katz 1994 377	104	NR	8	NR	NR	NR
Katz 1994 303	74	NR	8	NR	NR	NR
Levine 1993 ³⁹³	67	NR	13	NR	NR	NR

Results

Test-retest reliability

Two studies have reported that two tests, the CTS-I and the AMFHQ, give similar results when administered twice to the same subject. The correlation coefficients describing the test-retest reliability are shown in Table 152.

Table 152. Results of test-retest reliability tests

Study	Number of patients	Tests evaluated	Time between test administrations	Type of statistical comparison being made	Was the instrument reliable?
Alderson 1999 315	26	AMFHQ	NR	Intraclass correlation coefficient Reported to be consistent	Yes
Atroshi 1998 ³²⁶	22	CTS-I	24 hours	Correlation coefficient r = 0.71	Yes
Levine 1993 ³⁹³	67	CTS-I	24 hours	Pearson's correlation coefficient r = 0.93	Yes

Table 153. Results of response to treatment tests

Study	Number of patients	Test evaluated	Treatment	Time of testing months	Effect size hedges' d (95% CI) ^a	Was the instrument responsive to treatment?
Vaile 1999 475	27	mSHAQ	Injection of corticosteroids	<u>0</u>	0.31 (-0.23 to 0.85)	No
		SF-36	Injection of corticosteroids	0	-0.29 (-0.82 to 0.24)	No
		NHP	Injection of corticosteroids	<u>0</u> 1	0.38 (-0.16 to 0.91)	No
		V-VAS	Injection of corticosteroids	0	1.58 (0.97 to 2.19)	Yes
Atroshi 1998 ³²⁶	102	CTS-I	Carpal tunnel release surgery	0	0.78 (0.50 to 1.07)	Yes
	48	SF-36	Carpal tunnel release surgery	0	-0.052 (-0.45 to 0.35)	No
Katz 1994 303	43	CTS-I	Carpal tunnel release surgery	0	1.08 (0.63 to 1.53)	Yes
	55	K-ADL	Carpal tunnel release surgery	0 3	1.32 (0.91 to 1.73)	Yes
Levine 1993 ³⁹³	38	CTS-I	Carpal tunnel release surgery	0 14 mean	0.97 (0.50 to 1.45)	Yes

a calculated by ECRI

Validity

The validity tests performed on the instruments evaluated are summarized in Table 154. The validity tests can be separated into two groups: those measuring predictive validity, and those measuring concurrent validity.

Predictive validity

Atroshi 1997 compared the test scores of those receiving Workers' Compensation to the scores of those not receiving Workers' Compensation. Atroshi 1997 found no statistically significant differences between the two groups in their scores on either the SF-36 or the CTS-I. Workers' Compensation is paid to only those with injuries so severe that they cannot work. Thus, the results of this study suggest that either the SF-36 and the CTS-I are not valid tests for functional limitations, or that Workers' Compensation is not a valid measure of the severity of functional limitations. Due to a lack of reported data, we were unable to verify that the study by Atroshi 1997 had sufficient statistical power to be able to detect a statistical significance between the two groups if one had existed.

Table 154. Results of validity tests

Study	Number of patients	Test evaluated	Type of statistical comparison being made	Validated against	Was the instrument valid by this measurement?
Alderson 1999 ³¹⁵	26	AMHFQ	Pearson's correlation coefficient	pinch strength $r = 0.295$ grip strength $r = 0.3867$ two-point discrimination $r = -0.127$	Yes, but the r value is low Yes, but the r value is low No
Atroshi 1997 477	102	SF-36	ANOVA	On workers comp. vs. not on workers comp. p = 0.5	No
		CTS-I	ANOVA	On workers comp. vs. not on workers comp p = 0.07	No
Pransky 1997 ⁴⁷⁶	165	UEF	Difference between two means with t test	working vs. not working p <0.001 normal Phalen's test vs. abnormal Phalen's test p <0.05	Yes
			Pearson's correlation coefficient	nerve conduction speed test p >0.05 pinch strength p <0.001 grip strength p <0.001	No Yes Yes
Katz 1994 ³⁷⁷	104	Global score	Pearson's correlation coefficient	time to return to work- treated with open release surgery r = 0.67 time to return to work- treated with	Yes, but the r value is low
Levine 1993	67	CTS-I	Spearmann's	endoscopic release surgery r = 0.2 Semmes-Weinstein	Yes, but the r value is
393			correlation coefficient	monofilament testing $r = 0.24$ two-point discrimination test $r = 0.42$ pinch strength $r = 0.60$ grip strength $r = 0.50$ median nerve sensory conduction velocity $r = 0.12$	Yes Yes Yes No

Test-test comparisons

One study compared the scores of the same patients on different tests (Table 155). Atroshi 1998 compared the CTS-I and the SF-36 tests on patients with carpal tunnel syndrome. Before treatment of the carpal tunnel syndrome, the test scores correlated fairly well, but the correlation dropped after treatment. This change may be attributed to the finding, discussed previously, that the CTS-I instrument is responsive to treatment while the SF-36 is not.

Table 155. Results of test-test comparisons

Study	Tests being compared	Type of statistical comparison being made	Value of comparison r	Were the tests consistent?
Atroshi 1998 326	CTS-I and SF-36, pre-treatment	Spearmann's correlation coefficient	0.62	Yes
	CTS-I and SF-36, post-treatment	Spearmann's correlation coefficient	0.56	Yes

Conclusion

Eight studies evaluated the ability of nine different instruments as ways to measure functional limitations of patients with carpal tunnel syndrome. Of the available instruments, only two were evaluated by more than one trial. The two instruments that were evaluated by three and four trials, respectively, were the SF-36 and the Levine CTS-I.

It can be tentatively concluded that the SF-36 is not a useful instrument for assessing functional limitations in individuals with carpal tunnel syndrome. The SF-36 was reported to not be responsive to treatment and to not be able to predict ability to work.

It can be tentatively concluded that the Levine CTS-I may be a useful instrument for assessing functional limitations in individuals with carpal tunnel syndrome. This instrument was reported to be responsive to treatment, and to have concurrent validity as measured by grip and pinch strength. However, the Levine CTS-I was not evaluated by the studies included in the answer to this question for internal reliability, or prediction of the ability to perform activities of daily living. In addition, the Levine CTS-I has been reported by one study to not be able to predict ability to work.

It is difficult to reach an evidence-based conclusion as to the usefulness of the other instruments evaluated in this report due to the limited evidence base.

Question #10: What are the functional limitations for an individual with carpal tunnel syndrome before treatment?

This question inquires about the functional limitations of an individual before they have received conservative or surgical treatment for carpal tunnel syndrome. In addressing it, our objective is to catalogue these limitations, and not to address the effectiveness of these treatments. We address the effectiveness of conservative and surgical treatments in Question 3.

The available literature governs our approach to the present question. Hence, we address functional status rather than functional limitations, because no published studies specifically addressed the latter. In addition, the only available data operationally defines functional status in terms of scores on certain written tests. Hence, we also address functional status in these terms. The validity and reliability of these written tests is discussed in Question 9. Study inclusion criteria are described under Methods (section).

Excluded studies

As discussed in the Methods section, we retrieved articles identified by our literature searches according to certain *a priori* criteria. However, not all of the retrieved studies met our more specific inclusion criteria for this question. These latter studies, and the reason we did not consider them for this question are shown in Table 156.

Table 156. Excluded studies

Author	Reason for exclusion
Sefcovic	Some patients had prior treatment (including surgery), some
(2000) 481	did not, but all were analyzed together.
Davis (1998)	Used CTOA-I scale that has not been validated against
438	accepted functional scales for carpal tunnel syndrome

There were also nine studies wherein functional status was reported for patients prior to receiving surgical treatment. 44,311,313,326,428,476,482-484 These patients generally had received prior conservative treatment that had been ineffective at relieving symptoms (or had not provided enough relief). Because patients who eventually receive surgery may have more severe pre-treatment symptoms than non-surgical patients, these nine studies do not address the question and are not considered further.

Evidence Base

Two studies (with a total of 51 patients) remained that addressed this question after the above exclusions.

Internal validity

Aspects of study quality that are most relevant to the present question are shown in Table 157. Because we are cataloging functional status rather than using it to compare treatments, randomization and the use of control groups are not of paramount importance here. Therefore, Table 157 does not depict these aspects of study design. However, the following variables are particularly important: attrition rates, whether the trial was prospective, and whether the raters of functional status (in this case the patients) were blinded to the treatment the patient received.

One study reported no patient attrition, the other reported an attrition rate of 19 percent. This latter study did not perform an intent-to-treat analysis.³⁴ Both studies were prospective, but neither employed blinding. Because it is difficult to blind patients to the treatment received, we are considering unblinded studies to be of acceptable quality for this question.

Table 157. Internal validity

Author	Number of patients	Number of centers	Funded by a for-profit agency?	Prospective	Blinding	% Attrition	Intent to treat analysis	% Compliance
Walker (2000) 34	21	1	No	Yes	No	19.0	No	92
Vaile (1999) ⁴⁸⁵	30	2	NR	Yes	No	0	Yes	NR

NR – Not reported

Generalizability

Selected patient characteristics are presented in Table 158. Both studies reported mean patient age and percentage of female patients. For the remaining categories, one study reported combidities, ⁴⁸⁵ and neither study reported duration of symptoms or selection of patients based on severity of disease. In one study (Walker et al., 2000), the percent of female patients was much lower than that found in a typical population of carpal tunnel patients. This study examined a population of Veteran's Administration patients, of which men comprise an overwhelming majority. ³⁴ Although Vaile et al. (1999) did not report a mean age, the range suggests that the mean age is probably consistent with epidemiologic studies (see Introduction section, carpal tunnel syndrome, subheading epidemiology, as well as Question two for CTS).

Only one study reported any information relating to patient employment or occupation. Vaile et al. (1999) reported that there were no patients receiving workers' compensation (Table 159). Because there were only two studies, and they incompletely presented information on occupation-related variables, one cannot determine how generalizable these studies are to the greater population of patients with carpal tunnel syndrome.

Table 158. Patient characteristics

Author	Number of patients	Mean age (range)	% female	Duration of condition mean and range (months)	% Patients with diabetes	% Patients with arthritis	% Patients with prevous relevant injuries	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Did the study exclude patients with severe disease?	Did the study exclude patients with mild disease?
Walker (2000) 34	21	60 (44- 81)	4.8	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
Vaile (1999) 485	30	(29-84)	81.5	NR	NR	55.6	NR	7.4	NR	NR	NR	NR	NR

Table 159. Patient occupation

Author	Number of patients	% Patients employed	% Patients on Workers Compensation	% Patients retired	% Patients homemakers	Reported Occupations
Walker (2000) 34	21	NR	NR	NR	NR	NR
Vaile (1999) ⁴⁸⁵	30	NR	0	NR	NR	NR

Results

Table 160 shows the reported functional status of patients with carpal tunnel syndrome who had no prior treatment. Since each study used a different scale to measure functional status, the scores are not directly comparable. The two studies suggested that untreated patients on average score in the middle range (the 30-65% level) of functional status scales, suggesting mild to moderate difficulty with functional activities. 34,485

Table 160. Studies with patients who had no prior treatment

Study	N	Future treatment	Scale	Range of scale	Overall mean pre-treatment functional status score	% of maximum score
Walker (2000) ³⁴	21	Non-surgical (splints)	CTS-I	1-5	Splint (night only): 2.75 (1.01)	43.8
					Splint (full-time): 2.27 (1.03)	31.8
Vaile (1999) ⁴⁸⁵	30	Non-surgical (steroid injections)	Vaile VAS	0-100	64.2 (24.0)	64.2

CTS-I – Carpal Tunnel Syndrome Instrument

VAS – Visual Analog Scale

Conclusions

There is some evidence to suggest that most untreated patients with carpal tunnel syndrome have mild to moderate functional difficulties before treatment. However, this evidence is derived from only two studies comprised of a total of 51 patients. This is too few patients and too few studies to allow one to reach a firm evidence-based conclusion.

Question #11: What are the functional limitations of an individual with carpal tunnel syndrome after treatment?

This question inquires about the functional limitations of an individual after they have received conservative or surgical treatment for carpal tunnel syndrome. Our objective is as described in Question 10 for carpal tunnel syndrome. As also discussed in Question 10, our approach is governed by the available literature. We refer the reader to that question for additional details.

Excluded studies

As discussed in the Methods section, we retrieved articles identified by our literature searches according to certain *a priori* criteria. However, not all of the retrieved studies met our more specific inclusion criteria for this question. Table 161 shows these latter studies and the reason we did not consider them for this question.

Table 161. Excluded studies

Author	Reason for exclusion
Provinciali (2000) 427	Used Jebsen-Taylor test to measure functional limitation. The test is not validated for carpal tunnel syndrome
Atroshi (1999) 486	Study group overlaps with Atroshi et al. 326
Bessette (1998) ⁴⁸⁷	Used SF-36 scale that is not accurate for carpal tunnel syndrome (see Question 9 for carpal tunnel syndrome)
Davis (1998) 438	Used CTOA-I scale that has not been validated against accepted functional scales for carpal tunnel syndrome
Katz (1998) ⁴⁶²	Study group contains an unspecified number of the same patients evaluated in Katz et al. 482
Atroshi (1997) 483	Lack of information about treatment status of the study group
Katz (1996) ⁴⁸⁸	Study group contains an unspecified number of the same patients evaluated in Katz et al. ⁴⁸²
Katz (1994) 303	Biased post-hoc selection of patients for analysis

Evidence base

Twelve studies (with a total of 1567 patients) that addressed this question remained after the above exclusions.

Internal Validity

Aspects of study quality that are most relevant to the present question are shown in Table 162. Because we are cataloging functional status rather than using it to compare treatment, randomization and the use of control groups are not of paramount importance here. Therefore, Table 162 does not depict these aspects of study design. However, the

Table 162. Study quality

Author								
	Number of patients	Number of centers	Funded by a for-profit agency?	Prospective	Blinding	% Attrition	Intent to treat analysis	% Compliance
Mondelli (2000) 311	110	1	No	NR	No	15.5	No	NA
Porras (2000) 313	85	1	NR	Yes	No	0	Yes	NA
Walker (2000) 34	21	1	No	Yes	No	19.0	No	92
Vaile (1999) ⁴⁸⁵	30	2	NR	Yes	No	0	Yes	NR
Atroshi (1998) ³²⁶	111	1	No	Yes	No	8.1	No	NA
Katz (1998) ⁴⁸²	429	26	No	Yes	No	21 (6 months) 28 (18 months) 31 (30 months)	No	NR
Atroshi (1997) ⁴⁷⁷	277	1	No	NA	NA	24	No	NR
Pransky (1997) ⁴⁷⁶	165	1	No	Yes	No	13 37 (18 months)	No	NR
Amadio (1996) ⁴⁸⁴	22	1	No	Yes	No	0	Yes	NA
Blair (1996) ⁴²⁸	86	1	No	Yes	Single (partly)	11.8	No	NA
Worseg (1996) ⁴⁴	126	1	No	Yes	No	0	Yes	NA
Levine (1993) 393	105	1	No	Yes (partly)	No	Not clear	Yes	NR

Generalizability

Selected patient characteristics are presented in Table 163. Ten of 12 studies (83.3%) reported mean patient age and all studies reported percentage of female patients. The mean ages of patients in surgical studies (53.4 years) was similar to that reported in epidemiological studies (see Introduction section, subheading epidemiology) and the average obtained from the 124 surgical studies (50.5 years) that were evaluated for any question in this document (see Question 2). The percentage of female patients in surgical studies was generally similar to that observed when compared to all surgical studies. The

Table 163. Patient characteristics

Author	Number of patients	Mean age (range)	% female	Duration of condition mean and range (months)	% Patients with diabetes	% Patients with arthritis	% Patients with prevous	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Did the study exclude patients with severe disease?	Did the study exclude patients with mild disease?
Mondelli (2000) ³¹¹	110	56 (20-82)	86.0	NR	5.4	0	4.3	NR	1.1	NR	0	NR	NR
Porras (2000) 313	85	52 (18-81)	90.6	39 (6-300)	NR	NR	NR	NR	NR	NR	NR	NR	NR
Walker (2000)	21	60 (44-81)	4.8	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
Vaile (1999) 485	30	(29-84)	81.5	NR	NR	55.6	NR	7.4	NR	NR	NR	NR	NR
Atroshi (1998) ³²⁶	111	52 (21-88)	65.7	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
Katz (1998)	429	NR	74.2	NR	NR	NR	NR	NR	NR	0	NR	NR	NR
Atroshi (1997) ⁴⁷⁷	277	WC: 41 (25-62) Non-WC: 49 (13- 91)	77.8	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
Pransky (1997) ⁴⁷⁶	165	46 (22-80)	67	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
Amadio (1996) ⁴⁸⁴	22	60 (33-80)	59.1	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR

Author	Number of patients	mean age (range)	% fem	Duration of condition mean and range (months)	% Patients with diabetes	% Patients with arthritis	% Patients with prevous	% Patients with other relevant nerve impingement conditions	% Patients with peripheral neuropathy	% Patients pregnant	% Patients on kidney dialysis	Did the study exclude patients with severe disease?	Did the study exclude patients with mild disease?
Blair (1996) 428	86	49 (23-82)	82.7	NR	NR	NR	NR	NR	0	NR	NR	NR	NR
Worseg (1996) ⁴⁴	126	56 (35-90)	69.8	23.4	NR	NR	NR	NR	0	NR	NR	NR	NR
Levine (1993)	105	58 (19-88)	74.3	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR

Table 164. Patient occupation

Author	Number of patients	% Patients employed	% Patients on Workers Compensation	% Patients retired	% Patients homemakers	Reported Occupations
Mondelli (2000) 311	110	NR	NR	NR	NR	NR
Porras (2000) ³¹³	85	NR	NR	NR	NR	Homemaker, low functional demand, cleaners, keyboard workers, heavy work, assembly line
Walker (2000) 34	21	NR	NR	NR	NR	NR
Vaile (1999) 485	30	NR	0	NR	NR	NR
Atroshi (1998) 326	111	NR	NR	NR	NR	NR
Katz (1998) 482	429	NR	38.2	NR	NR	NR
Atroshi (1997) 477	277	NR	28.8	NR	NR	NR
Pransky (1997) 476	165	89	10	NR	NR	NR
Amadio (1996) 484	22	63.6	0.9	NR	NR	NR
Blair (1996) 428	86	NR	NR	NR	NR	NR
Worseg (1996) 44	126	NR	NR	47.6	6.3	Retired, employee, worker, unemployed, homemaker, other
Levine (1993) 393	105	NR	12.4	NR	NR	NR

Results

Table 165 shows the results of the two nonsurgical studies of post-treatment functional limitations in patients with carpal tunnel syndrome. Since these studies used different scales to measure functional status, their scores are not directly comparable. Both studies suggested that after nonsurgical treatment, patients score, on average, in the lower range (the 20-30% level) of functional status scales. However, it is unclear whether the results of these two studies are generalizable to the larger patient population.

Table 165. Studies with patients who had no prior treatment

Study	N	Treatment	Scale	Range of scale	Overall mean post-treatment functional status score (± SD)	% of maximum score
Walker (2000) ³⁴	21	Non-surgical (splints)	CTS-I	1-5	Splint (night only): 2.14 (0.87) Splint (full-time): 1.93 (0.77)	28.5 23.3
Vaile (1999) ⁴⁸⁵	30	Non-surgical (steroid injections)	Vaile VAS	0-100	23.8 (26.2)	23.8

CTS-I – Carpal Tunnel Syndrome Instrument

VAS - Visual Analog Scale

Table 166 shows the results of the two surgical studies that reported individual functional activity mean scores using the CTS-I scale. Lower scores on this scale indicate less functional limitation. Table 167 shows the number of patients for each level of the scale in the surgical study of Atroshi et al. (1998). 326

Table 168 shows the results of a third surgical study, performed by Blair et al. 428 Although these latter authors did not use a specific scale, they did report the number of patients who had difficulty with specific functional activities. Both of these studies suggest that patients have relatively mild functional limitations following surgery, and the study by Blair et al. suggests that the majority of patients do not have any noticeable difficulty with certain functional activities after surgery.

Seven studies reported overall mean functional activity scores on the CTS-I scale prior to surgery (Table 169). Four out of seven studies did not describe the surgical procedure, so no evidence-based conclusions can be reached concerning functional limitations after specific surgical procedures. However, one can make some broad conclusions about functional limitations after surgical procedures as a group. These studies suggested that most patients report no-to-moderate difficulty with functional activities (mean 1.4-2.6 on CTS-I) after surgery. Although there were no statistically significant posttreatment differences between specific patient groups, in two studies there was a trend toward more difficulty with functional activities among patients receiving workers' compensation.

Table 166. Studies with post-treatment functional limitation data for patients with carpal tunnel syndrome (individual functional activities – mean scores from CTS-I)

Study	N	Treatment	Range of scale	Writing	Holding a book	Buttoning clothes	Gripping the telephone	Opening jars	Performing household chores	Carrying a grocery bag	Bathing and dressing
Atroshi (1998) ³²⁶	111	Endoscopic release	1-5	1.5	1.7	1.7	1.5	2.1	1.7	2.1	1.3
Worseg (1996) ⁴⁴	126	Endoscopic release	1-5	1.0 (0.2)a	1.0 (0.1)	1.0 (0.1)	1.0 (0.1)	1.6 (0.7)	1.4 (0.8)	1.4 (0.8)	1.2 (0.4)
		Open release		1.0 (0.2)	1.0 (0.2)	1.2 (0.4)	1.1 (0.2)	1.9 (0.8)	1.2 (0.4)	1.7 (0.8)	1.2 (0.4)

^aNumbers in parentheses indicate standard deviations

Table 167. Studies with post-treatment functional limitation data for patients with carpal tunnel syndrome (individual functional activities – number of patients)

Study	N	Score	Number	lumber of patients in each CTS-I Functional Status category (%)								
			Writing	Writing Holding a book Buttoning Gripping the clothes Clothes Gripping the jars		Opening jars	Performing household chores	Carrying a grocery bag	Bathing and dressing			
Atroshi	111	1	69 (70.4)	59 (60.2)	59 (59.6)	69 (72.6)	42 (42.4)	56 (56.6)	41 (42.3)	77 (77)		
(1998) ³²⁶		2	17 (17.3)	21 (21.4)	19 (19.2)	12 (12.6)	26 (26.3)	21 (21.2)	25 (25.8)	18 (18)		
		3	6 (6.1)	9 (9.2)	15 (15.2)	7 (7.4)	13 (13.1)	16 (16.2)	16 (16.5)	3 (3)		
		4	6(6.1)	9 (9.2)	2 (2.0)	4 (4.2)	14 (14.1)	4 (4.0)	12 (12. 4)	2 (2)		
		5	0 (0)	0 (0)	4 (4.0)	3 (3.2)	4 (4.0)	1 (1.0)	3 (3.1)	0 (0)		

Table 168. Studies with post-treatment functional limitation data for patients with carpal tunnel syndrome (individual functional activities – number of patients)

Study	Treatment	Difficulty	Self-described difficulty in performing selected activities of daily living after carpal tunnel release (% of patients)			
			Screwing lids	Picking up small objects	Lifting	
Blair (1996) ⁴²⁸	Open release plus epineurotomy (n = 48)	Yes No	15 (31.3) 33 (68.8)	9 (18.8) 39 (81.3)	9 (18.8) 39 (81.3)	
	Open release without epineurotomy (n = 27)	Yes No	11 (40.7) 16 (59.3)	10 (37.0) 17 (63.0)	7 (25.9) 20 (74.1)	

Table 169. Studies with post-treatment functional limitation data for patients with carpal tunnel syndrome (mean function score on CTS-I)

Study	N	Treatment	Study Design	Range of scale	Followup time	Overall mean post- treatment functional status score (SD)	% of maximum score
Mondelli (2000) 311	110	Surgical (open release)	Prospective case series	1-5	1 month 6 months	2.0 (0.7)	25 12.5
Porras (2000) 313	85	Surgical (open release)	Prospective case series	1-5	6 months	1.4 (range 1- 4.2)	10
Atroshi (1998) ³²⁶	111	Surgical (endoscopic release)	Prospective case series	1-5	3 months	1.7 (range 1.6- 1.9)	17.5
Katz (1998) ⁴⁸²	429	Surgical (n = 270, procedures not described)	Prospective case series (stratified)	1-5	6 months	Surgical patients: >55 years: 1.7 (0.9)	17.5
		Non-surgical (n = 125) (34 patients				≤55 years, WC non-recipient:: 1.6 (0.7)	15
		who crossed over to surgery were				≤55 years, WC recipient: 2.1 (0.9)	27.5
		not evaluated)				Non-surgical patients: >55 years: 2.6 (0.8)	40
						≤55 years, WC non-recipient:: 1.9 (0.9)	22.5
						≤55 years, WC recipient: 2.2 (0.7)	30
					18 months	Surgical patients: >55 years: 1.6 (0.7)	15
						≤55 years, WC non-recipient: 1.6 (0.7)	15
						≤55 years, WC recipient: 2.2 (0.9)	30

Study	N	Treatment	Study Design	Range of scale	Followup time	Overall mean post- treatment	% of maximum score
				Scale		functional	Score
						status	
						score (SD)	
						Non-surgical patients:	
						>55 years:	32.5
						2.3 (0.9)	
						≤55 years, WC	
						non-recipient::	25
						2.0 (1.0)	25
						≤55 years,	0.5
						WC recipient:	35
					30 months	2.4 (0.7)	
					30 1110111115	Surgical patients:	
						>55 years:	15
						1.6 (0.9)	
						≤55 years, WC	
						non-recipient:	15
						1.6 (0.7)	15
						≤55 years, WC	
						recipient:	30
						2.2 (1.0) Non-surgical	
						patients:	
						>55 years:	30
						2.2 (0.8)	
						≤55 years, WC	
						non-recipient::	
						2.0 (0.9)	25
						≤55 years,	20
						WC recipient: 2.2 (0.8)	30
Atroshi	277	Surgical or	Cross-	1-5	6-20 months	WC patients:	
(1997) 477		non-surgical	sectional		3 20	2.5 (95% CI:	37.5
		(or both)	study			2.2-2.7)	
		(procedures not described)				Non-WC	
		not described)				patients:	30
						2.2 (2.0-2.4)	
Amadio	22	Surgical (not	Prospective	1-5	3 months	1.77 (0.68)	19.3
(1996) 484		described)	case series				

Study	N	Treatment	Study Design	Range of scale	Followup time	Overall mean post- treatment functional status score (SD)	% of maximum score
Levine (1993) 393	67	Surgical or non-surgical (not described)	Prospective case series	1-5	3 months	Prospective: 2.1 (1.1)	27.5
	38	Surgical (not described)	Retrospective case series		Median: 14 months	Retrospective: 2.0 (1.1)	25

WC – Workers' Compensation

Table 170. Studies with post-treatment functional limitation data for patients with carpal tunnel syndrome (summary function score on UEFS)

Study	N	Treatment	Study Design	Range of scale	Followup time	Overall summary post-treatment functional status score (SD)	% of maximum score
Pransky (1997) ⁴⁷⁶	108	Surgical or non-surgical (not described)	Prospective case series	1-10	Mean: 18 months	25.4 (18.1)* Note: this study also had a case series of mixed upper extremity disorders (UEDs)	17.1

Conclusions

Although studies of non-surgical therapies suggested that most patients experience only mild difficulty with functional activities after treatment, it is unclear whether the results of these two studies are generalizable to the larger patient population. Studies with surgical outcomes suggested that most patients report no-to-moderate difficulty with functional activities (mean 1.4-2.6 on CTS-I) after surgery. Although there were no statistically significant differences between specific patient groups, in two studies there was a trend toward more difficulty with functional activities among workers' compensation patients. Decreased functional ability on the CTS-I scale shows a strong correlation with work absence. The available data are insufficient to determine a cutoff point on measuring scales above which patients are unable to work.